

IN THE UNITED STATES DISTRICT COURT
FOR THE EASTERN DISTRICT OF PENNSYLVANIA

MALLINCKRODT IP and MALLINCKRODT)
HOSPITAL PRODUCTS INC.,)
)
Plaintiffs,)
)
v.) C.A. No. _____
)
B. BRAUN MEDICAL INC.) **COMPLAINT FOR PATENT**
) **INFRINGEMENT**
)
Defendant.)

COMPLAINT FOR PATENT INFRINGEMENT

Plaintiffs Mallinckrodt IP and Mallinckrodt Hospital Products, Inc. (“Plaintiffs”) for their Complaint against defendant B. Braun Medical Inc. (“Braun”), allege as follows:

PARTIES

1. Plaintiff Mallinckrodt IP is a company organized and existing under the laws of Ireland, having a registered address of Damastown Industrial Estate, Mulhaddart, Dublin 15, Ireland. Mallinckrodt IP is a wholly-owned subsidiary of Mallinckrodt plc. As set forth herein, Mallinckrodt IP is the assignee of U.S. Patent No. 9,399,012 (“the ’012 patent”) and U.S. Patent No. 9,610,265 (“the ’265 patent”) (collectively, the “patents-in-suit”).

2. Plaintiff Mallinckrodt Hospital Products Inc. (“Mallinckrodt Hospital Products”), formerly Cadence Pharmaceuticals, Inc. (“Cadence”), is a company organized and existing under the laws of Delaware, having a principal place of business at 675 McDonnell Blvd., Hazelwood, Missouri 63042. Mallinckrodt Hospital Products is a wholly-owned subsidiary of Mallinckrodt plc.

3. Upon information and belief, Defendant Braun is a company organized under the

laws of Pennsylvania, having a principal place of business at 824 Twelfth Avenue, Bethlehem, Pennsylvania 18018. Upon information and belief, Braun is in the business of manufacturing, distributing, and selling pharmaceutical products throughout the United States, including in this judicial district.

NATURE OF THE ACTION

4. This is a civil action for infringement of the patents-in-suit pursuant to the Patent Laws of the United States, 35 U.S.C. §§ 100 *et seq.*; the Federal Food, Drug, and Cosmetic Act; and the Declaratory Judgment Act, 28 U.S.C. §§ 2201 *et seq.*

JURISDICTION AND VENUE

5. This Court has subject matter jurisdiction over this action under 28 U.S.C. §§ 1331, 1338(a), and 2201(a).

6. This Court has personal jurisdiction over Braun because, upon information and belief, *inter alia*, Braun is Pennsylvania company and has a principal place of business in this District. This Court has personal jurisdiction over Braun for the additional reasons set forth below and for other reasons that will be presented to the Court if such jurisdiction is challenged.

7. This Court has personal jurisdiction over Braun because, *inter alia*, Braun has purposefully availed itself of the rights and benefits of Pennsylvania law by engaging in systematic and continuous contacts with Pennsylvania.

8. Upon information and belief, Braun regularly and continuously transacts business within the Commonwealth of Pennsylvania, including by selling pharmaceutical products in Pennsylvania. Upon information and belief, Braun derives substantial revenue from the sale of those products in Pennsylvania and has availed itself of the privilege of conducting business within the Commonwealth of Pennsylvania. Upon information and belief, Braun regularly and continuously transacts business within the Commonwealth of Pennsylvania, including by selling

pharmaceutical products in Pennsylvania. Upon information and belief, Braun derives substantial revenue from the sale of those products in Pennsylvania and has availed itself of the privilege of conducting business within the Commonwealth of Pennsylvania. Upon information and belief, Braun provides maintenance and support services for its products in the Commonwealth of Pennsylvania.

9. On information and belief, Braun has purposefully availed itself of the benefits of this district by filing patent-related complaints in this district. *See B. Braun Med. Inc. v. Injectimed, Inc.*, 02-cv-09362; *B. Braun Med. Inc. v. Johnson & Johnson*, No. 00-cv-00380; *B. Braun Med. Inc. v. Becton, Dickinson & Co.*, No. 00-cv-00141.

10. This Court has personal jurisdiction over Braun because, *inter alia*, upon information and belief, Braun has submitted New Drug Application (“NDA”) No. 204957, claiming bioequivalence to Plaintiffs’ OFIRMEV® injectable acetaminophen product and seeking nationwide approval of its proposed product. Braun’s submission of NDA No. 204957 constitutes infringement of the patents-in-suit pursuant to 35 U.S.C. § 271(e). Braun’s tortious act of infringing the patents-in-suit causes concrete harm to Plaintiffs in the Commonwealth of Pennsylvania. By a letter received by Plaintiffs on February 23, 2017 (the “Braun Letter”), Braun stated that it had submitted NDA No. 204957 seeking approval to engage in the commercial manufacture, use, sale or offer for sale, and/or importation of acetaminophen for injection 10 mg/ml (“Braun’s Generic Product”) prior to the expiration of the ’218 patent. By a second letter received by Plaintiffs on April 21, 2017 (the “Second Braun Letter”), Braun stated that it had contemporaneously submitted an amendment to its NDA “to further indicate and confirm its intent” to engage in the commercial manufacture, use, sale or offer for sale, and/or importation of Braun’s Generic Product prior to the expiration of the ’012 patent.

11. Venue is proper in this Court pursuant to 28 U.S.C. § 1400(b). Under the Hatch-Waxman Act, the evaluation of infringement involves what the applicant will “likely market if its application is approved.” *Bayer AG v. Elan Pharm. Research Corp.*, 212 F.3d 1241, 1248-49 (Fed. Cir. 2000) (citing *Glaxo, Inc. v. Novopharm, Ltd.*, 110 F.3d 1562, 1569 (Fed. Cir. 1997)). Braun’s submission of NDA No. 204957, claiming bioequivalence to Plaintiffs’ OFIRMEV® injectable acetaminophen product and seeking nationwide approval of Braun’s Generic Product is an act of infringement of the patents-in-suit in this district pursuant to 35 U.S.C. § 271(e) causing concrete harm to Plaintiffs. Thus, in the context of the Hatch-Waxman Act, Braun has committed an act of infringement directed to and/or within this district.

12. Upon information and belief, Braun regularly and continuously transacts business within the Commonwealth of Pennsylvania, including by selling pharmaceutical products and medical equipment in Pennsylvania. Upon information and belief, Braun derives substantial revenue from the sale of those products in Pennsylvania. Upon information and belief, Braun provides maintenance and support services for its products in the Commonwealth of Pennsylvania. Furthermore, upon information and belief, Braun has a principal place of business as well as other facilities in this District. Thus, on information and belief, Braun resides in this District. Independently, Braun has a regular and established place of business in this District. Therefore, venue is proper in this judicial district under 28 U.S.C. § 1400(b).

13. The ’012 patent is currently at issue in the action captioned *Mallinckrodt IP v. B. Braun Medical Inc.*, No. 17-cv-01521, filed after Plaintiffs received the Braun Letter but before Plaintiffs received the Second Braun Letter. The ’012 patent is also currently at issue in the action captioned *Mallinckrodt IP v. B. Braun Medical Inc.*, No. 17-365 (D. Del.), filed after Plaintiffs received the Braun Letter but before Plaintiffs received the Second Braun Letter. Both

the '012 patent and the '265 patent are at issue in *Mallinckrodt IP v. B. Braun Medical Inc.*, No. 17-660 (D.Del.), filed after Plaintiffs received the Second Braun Letter.

THE PATENTS-IN-SUIT

14. The '012 patent, titled "Reduced Dose Intravenous Acetaminophen," was duly and legally issued by the PTO on July 26, 2016. The named inventors assigned the application that issued as the '012 patent to Cadence, which subsequently assigned that application to Mallinckrodt IP. Mallinckrodt IP is now the sole assignee of the '012 patent. A true and correct copy of the '012 patent is attached as Exhibit A.

15. Claim 1 of the '012 patent recites "[a] method for the treatment of pain or fever in an adult human or an adolescent human subject weighing at least 50 kg, in need thereof, comprising administering to the subject, by an intravenous route of administration, a therapeutically effective amount of a pharmaceutical composition comprising about 550 mg to about 800 mg of acetaminophen; and repeating said administration at least once at an interval of about 3 to about 5 hours."

16. Claim 39 of the '012 patent recites "[t]he method of claim 1, wherein the administered dose of acetaminophen is 650 mg, and further comprising repeating intravenous administration of 650 mg acetaminophen at least once at an interval of about 3 hours to about 5 hours."

17. The '265 patent, titled "Reduced Dose Intravenous Acetaminophen," was duly and legally issued by the PTO on April 4, 2017. The named inventors assigned the application that issued as the '265 patent to Cadence, which subsequently assigned that application to Mallinckrodt IP. Mallinckrodt IP is now the sole assignee of the '265 patent. A true and correct copy of the '265 patent is attached as Exhibit B.

18. Claim 1 of the '265 patent recites "[a] method of treating pain in a human subject

weighing at least 50 kg comprising: co-administering to the subject a therapeutically effective amount of a first pharmaceutical composition comprising about 500 mg to about 750 mg of acetaminophen and a therapeutically effective amount a second pharmaceutical composition comprising an opioid analgesic; wherein the first pharmaceutical composition is administered to the subject intravenously.”

19. Claim 7 of the '265 patent recites “[t]he method of claim 1, wherein the first pharmaceutical composition comprises about 650 mg of acetaminophen.”

OFIRMEV®

20. Cadence obtained approval from the Food and Drug Administration (the “FDA”) for NDA No. 022450 for OFIRMEV®, the first and only intravenous (IV) formulation of acetaminophen available in the United States. As part of the corporate restructuring resulting from the purchase of Cadence by Mallinckrodt plc, Mallinckrodt IP is now the holder of NDA No. 022450. Mallinckrodt Hospital Products distributes OFIRMEV®.

21. OFIRMEV® was approved by the FDA on November 2, 2010. OFIRMEV® is indicated for the management of mild to moderate pain, management of moderate to severe pain with adjunctive opioid analgesics, and reduction of fever.

22. The publication “Approved Drug Products with Therapeutic Equivalence Evaluations” (the “Orange Book”) identifies drug products approved on the basis of safety and effectiveness by the FDA under the Federal Food, Drug, and Cosmetic Act. Pursuant to 21 U.S.C. § 355(b)(1) and attendant FDA regulations, the '012, and '265 patents were timely listed in the Orange Book with respect to OFIRMEV®.

DEFENDANT'S INFRINGEMENT OF THE PATENTS-IN-SUIT

23. Upon information and belief, Braun submitted NDA No. 204957 to the FDA under the Federal Food, Drug, and Cosmetic Act (21 U.S.C. § 355(b)), seeking approval to

engage in the commercial manufacture, use, sale or offer for sale, and/or importation of Braun's Generic Product, prior to the expiration of the '012 patent and the '265 patent, both of which are listed in the Orange Book with respect to OFIRMEV®.

24. In the Second Braun Letter, Braun stated that it had submitted an amendment to NDA No. 204957 seeking approval to engage in the commercial manufacture, use, sale or offer for sale, and/or importation of Braun's Generic Product prior to the expiration of the '012 patent.

25. The Second Braun Letter also stated that the amendment to NDA No. 204957 contained a certification under 21 U.S.C. § 355(b)(2)(A)(iv) (the "Paragraph IV certification") alleging that the '012 patent is "invalid, unenforceable and/or will not be infringed by the manufacture, use, or sale of the product for [sic: which] B. Braun's NDA is submitted."

26. Braun's submission of NDA No. 204957 to the FDA, including its Paragraph IV certification, constitutes an act of infringement of the '012 patent under 35 U.S.C. § 271(e)(2)(A). In the event that Braun commercially manufactures, imports, uses, offers for sale, or sells Braun's Generic Product or induces such conduct, said actions would constitute infringement of the '012 patent under 35 U.S.C. § 271(a) and/or (b).

27. Upon information and belief, the FDA will require the labeling for Braun's Generic Product to be substantially identical to the approved labeling for OFIRMEV®, and Braun's Generic Product, if approved, will be marketed, sold, and/or distributed with labeling that is substantially identical to the labeling for OFIRMEV®.

28. The OFIRMEV® labeling includes instructions for administering OFIRMEV® to treat pain or fever in an adult human or an adolescent human subject weighing at least 50 kg, in need thereof, by administering to the subject, by an intravenous route of administration, a therapeutically effective amount of a pharmaceutical composition comprising 650 mg of

acetaminophen and repeating said administration at least once at an interval of 4 hours. A true and correct copy of the OFIRMEV® labeling is attached as Exhibit C.

29. For instance, Section 2.2 of the OFIRMEV® labeling recites that for adults and adolescents weighing 50 kg and over, “the recommended dosage of OFIRMEV is 1000 mg every 6 hours or 650 mg every 4 hours, with a maximum single dose of OFIRMEV of 1000 mg, a minimum dosing interval of 4 hours, and a maximum daily dose of acetaminophen of 4000 mg per day.”

30. Table 1 of the OFIRMEV® labeling also contains recommended dosing information for adults and adolescents weighing 50 kg and over, reciting that the “[d]ose given every 4 hours” is “650 mg.”

31. Section 2.4 of the OFIRMEV® labeling provides instructions and/or recommendations for dosing and recites, in pertinent part, that “[f]or doses less than 1000 mg, the appropriate dose must be withdrawn from the vial and placed into a separate container prior to administration. Using aseptic technique, withdraw the appropriate dose (650 mg or weight-based) from an intact sealed OFIRMEV vial and place the measured dose in a separate empty, sterile container (e.g. glass bottle, plastic intravenous container, or syringe) for intravenous infusion . . .”

32. Section 6.1 of the OFIRMEV® labeling reports on clinical trials in which patients were administered 650 mg OFIRMEV® every 4 hours.

33. Section 14.1 of the OFIRMEV® labeling describes acute pain studies in adults in which patients were administered 650 mg OFIRMEV® every 4 hours. The OFIRMEV® labeling reports that patients receiving OFIRMEV® experienced a statistically significant greater reduction in pain intensity over 24 hours compared to placebo.

34. The OFIRMEV® labeling therefore instructs, recommends, promotes, and/or encourages medical care providers to practice the methods of at least Claims 1 and 39 of the '012 patent.

35. The foregoing information in the OFIRMEV® labeling is essential for the safe and effective use of the drug, particularly given the warnings in the labeling concerning potential dosing errors. As the warning in the Highlights of Prescribing Information indicates, “[t]ake care when prescribing, preparing, and administering OFIRMEV Injection to avoid dosing errors which could result in accidental overdose and death.” The Highlights continue: “Acetaminophen has been associated with cases of acute liver failure, at times resulting in liver transplant and death. Most of the cases of liver injury are associated with the use of acetaminophen at doses that exceed the recommended maximum daily limits . . .”

36. Under the Hatch-Waxman Act, the evaluation of infringement involves what the applicant will “likely market if its application is approved.” *Bayer AG v. Elan Pharm. Research Corp.*, 212 F.3d 1241, 1248-49 (Fed. Cir. 2000) (citing *Glaxo, Inc. v. Novopharm, Ltd.*, 110 F.3d 1562, 1569 (Fed. Cir. 1997)).

37. Upon information and belief, the FDA will require the labeling for Braun's Generic Product, if approved, to contain recommendations and/or instructions that are identical or substantially identical to those set forth above from the OFIRMEV® labeling and, therefore, will contain recommendations and/or instructions for treating pain or fever in an adult human or an adolescent human subject weighing at least 50 kg, in need thereof, by administering to the subject, by an intravenous route of administration, a therapeutically effective amount of a pharmaceutical composition comprising 650 mg of acetaminophen and repeating said administration at least once at an interval of 4 hours.

38. Upon information and belief, based on the labeling that is likely to be required by the FDA for Braun's Generic Product, if approved, Braun's Generic Product will be administered to treat pain or fever in an adult human or an adolescent human subject weighing at least 50 kg, in need thereof, by administering to the subject, by an intravenous route of administration, a therapeutically effective amount of a pharmaceutical composition comprising 650 mg of acetaminophen and repeating said administration at least once at an interval of 4 hours, which administration will constitute direct infringement of at least Claims 1 and 39 of the '012 patent. Upon information and belief, this will occur at Defendant's active behest, and with Defendant's intent, knowledge, and encouragement. Upon information and belief, Defendant will actively induce, encourage, and abet this infringement with knowledge that it is in contravention of Plaintiffs' rights under the '012 patent.

39. Braun's submission of NDA No. 204957 to the FDA constitutes an act of infringement of the '012 patent under 35 USC § 271(e)(2)(A). Moreover, Braun intends to commercially manufacture, import, use, offer for sale, or sell Braun's Generic Product and/or induce such conduct. Said actions would constitute infringement of the '012 patent under 35 USC § 271(a) and/or (b).

40. Pursuant to 21 U.S.C. §§355(b)(2)-(3), because the '265 patent was timely listed in the Orange Book, Braun is obligated to provide a patent certification with respect to the '265 patent and to notify Plaintiffs of its certification. That listing occurred on or about May 4, 2017. Upon information and belief, Braun is required to submit a certification under 21 U.S.C. § 355(b)(2)(A)(iv) with regard to the '265 patent.

41. However, Plaintiffs have not received any certification from Braun with respect to the '265 patent. Pursuant to 35 U.S.C. § 271(e)(2), “[i]t shall be an act of infringement to submit

an application under section 505(j) of the Federal Food, Drug, and Cosmetic Act or described in section 505(b)(2) of such Act for a drug claimed in a patent or the use of which is claimed in a patent.” 35 U.S.C. § 271(e)(2). Braun’s submission of its NDA seeking approval for Braun’s Generic Product is an act of infringement with regard to one or more claims of the Orange Book-listed ’265 patent pursuant to 35 U.S.C. § 271(e)(2). *See, e.g., The Meds. Co. v. Eagle Pharm., Inc.*, No. 16-569, 2016 WL 4418230 (D.N.J. Aug. 17, 2016) (holding jurisdiction existed where patents asserted were not listed in Orange Book); *Research Found. of State Univ. of N.Y. v. Mylan Pharm. Inc.*, No. 09-184, 2012 WL 1901267, at *4 (D. Del. May 25, 2012) (“[A] Paragraph IV certification against the [patent] is not required [for a patentee] to bring suit under Section 271(e)(2)”).

42. Upon information and belief, the FDA will require the labeling for Braun’s Generic Product to be substantially identical to the approved labeling for OFIRMEV®, and Braun’s Generic Product, if approved, will be marketed, sold, and/or distributed with labeling that is substantially identical to the labeling for OFIRMEV®.

43. The OFIRMEV® labeling includes instructions for administering OFIRMEV® to treat pain in a human subject weighing at least 50 kg by co-administering to the subject a therapeutically effective amount of a first pharmaceutical composition comprising about 650 mg of acetaminophen and a therapeutically effective amount of a second pharmaceutical composition comprising an opioid analgesic, wherein the first pharmaceutical composition is administered to the subject intravenously.

44. For example, Section 1 of the OFIRMEV® labeling provides that “OFIRMEV (acetaminophen) injection is indicated for the . . . Management of moderate to severe pain with adjunctive opioid analgesics.”

45. Section 2.2 of the OFIRMEV® labeling recites that for adults and adolescents weighing 50 kg and over, “the recommended dosage of OFIRMEV is 1000 mg every 6 hours or 650 mg every 4 hours, with a maximum single dose of OFIRMEV of 1000 mg, a minimum dosing interval of 4 hours, and a maximum daily dose of acetaminophen of 4000 mg per day.”

46. Table 1 of the OFIRMEV® labeling also contains recommended dosing information for adults and adolescents weighing 50 kg and over, reciting that the “[d]ose given every 4 hours” is “650 mg.”

47. Section 2.4 of the OFIRMEV® labeling provides instructions and/or recommendations for dosing and recites, in pertinent part, that “[f]or doses less than 1000 mg, the appropriate dose must be withdrawn from the vial and placed into a separate container prior to administration. Using aseptic technique, withdraw the appropriate dose (650 mg or weight-based) from an intact sealed OFIRMEV vial and place the measured dose in a separate empty, sterile container (e.g. glass bottle, plastic intravenous container, or syringe) for intravenous infusion . . .”

48. Section 6.1 of the OFIRMEV® labeling reports on clinical trials in which patients were administered 650 mg OFIRMEV® every 4 hours.

49. Section 14.1 of the OFIRMEV® labeling describes acute pain studies in adults in which patients were administered 650 mg OFIRMEV® every 4 hours. The OFIRMEV® labeling reports that patients receiving OFIRMEV® experienced a statistically significant greater reduction in pain intensity over 24 hours compared to placebo.

50. The OFIRMEV® labeling therefore instructs, recommends, promotes, and/or encourages medical care providers to practice the methods of at least Claim 7 of the '265 patent.

51. The foregoing information in the OFIRMEV® labeling is essential for the safe and effective use of the drug, particularly given the warnings in the labeling concerning potential dosing errors.

52. Under the Hatch-Waxman Act, the evaluation of infringement involves what the applicant will “likely market if its application is approved.” *Bayer AG v. Elan Pharm. Research Corp.*, 212 F.3d 1241, 1248-49 (Fed. Cir. 2000) (citing *Glaxo, Inc. v. Novopharm, Ltd.*, 110 F.3d 1562, 1569 (Fed. Cir. 1997)).

53. Upon information and belief, the FDA will require the labeling for Braun’s Generic Product, if approved, to contain recommendations and/or instructions that are identical or substantially identical to those set forth above from the OFIRMEV® labeling and, therefore, will contain recommendations and/or instructions for treating pain in a human subject weighing at least 50 kg by co-administering to the subject a therapeutically effective amount of a first pharmaceutical composition comprising about 650 mg of acetaminophen and a therapeutically effective amount of a second pharmaceutical composition comprising an opioid analgesic, wherein the first pharmaceutical composition is administered to the subject intravenously.

54. Upon information and belief, based on the labeling that is likely to be required by the FDA for Braun’s Generic Product, if approved, Braun’s Generic Product will be administered to treat pain in a human subject weighing at least 50 kg by co-administering to the subject a therapeutically effective amount of a first pharmaceutical composition comprising about 650 mg of acetaminophen and a therapeutically effective amount of a second pharmaceutical composition comprising an opioid analgesic, wherein the first pharmaceutical composition is administered to the subject intravenously, which co-administration will constitute direct infringement of at least Claim 7 of the ’265 patent. Upon information and belief, this will

occur at Defendant's active behest, and with Defendant's intent, knowledge, and encouragement. Upon information and belief, Defendant will actively induce, encourage, and abet this infringement with knowledge that it is in contravention of Plaintiffs' rights under the '265 patent.

55. Braun's submission of NDA No. 204957 to the FDA constitutes an act of infringement of the '265 patent under 35 USC § 271(e)(2)(A). Moreover, Braun intends to commercially manufacture, import, use, offer for sale, or sell Braun's Generic Product and/or induce such conduct. Said actions would constitute infringement of the '265 patent under 35 USC § 271(a) and/or (b).

56. Upon information and belief, Braun was aware of the '012 patent prior to filing NDA No. 204957, and its actions render this an exceptional case under 35 U.S.C. § 285.

57. Upon information and belief, Braun was aware of the application that subsequently issued as the '265 patent while continuing to seek approval of NDA No. 204957, and their actions render this an exceptional case under 35 U.S.C. § 285.

58. The acts of infringement by Defendant set forth above will cause Plaintiffs irreparable harm for which they has no adequate remedy at law, and will continue unless enjoined by this Court.

COUNT I
(INFRINGEMENT OF THE '012 PATENT)

59. Plaintiffs incorporate each of the preceding paragraphs 1 to 58 as if fully set forth herein.

60. Braun's submission of NDA No. 204957, including its Paragraph IV certification, constitutes infringement of the '012 patent pursuant to 35 U.S.C. § 271(e)(2).

61. Upon information and belief, upon FDA approval of NDA No. 204957, Braun will induce infringement of at least Claims 1 and 39 of the '012 patent by making, using,

offering to sell, or selling Braun's Generic Product in the United States, and/or importing Braun's Generic Product into the United States, in violation of 35 U.S.C. § 271.

62. Upon information and belief, upon FDA approval of NDA No. 204957, doctors, nurses, and other medical professionals will directly infringe at least Claims 1 and 39 of the '012 patent by using Braun's Generic Product, in violation of 35 U.S.C. § 271(a). Braun's Generic Product will be administered to treat pain or fever in an adult human or an adolescent human subject weighing at least 50 kg, in need thereof, by administering to the subject, by an intravenous route of administration, a therapeutically effective amount of a pharmaceutical composition comprising 650 mg of acetaminophen and repeating said administration at least once at an interval of 4 hours, which administration will constitute direct infringement of at least Claims 1 and 39 of the '012 patent.

63. Upon information and belief, this direct infringement will occur at Braun's active behest, and with Braun's intent, knowledge, and encouragement. Braun will intentionally encourage infringement of at least Claims 1 and 39 of the '012 patent at least by way of the labeling for Braun's Generic Product which will contain recommendations and/or instructions for treating pain or fever in an adult human or an adolescent human subject weighing at least 50 kg, in need thereof, by administering to the subject, by an intravenous route of administration, a therapeutically effective amount of a pharmaceutical composition comprising 650 mg of acetaminophen and repeating said administration at least once at an interval of 4 hours.

64. Upon information and belief, Braun is aware of the '012 patent, which is listed in the Orange Book with respect to OFIRMEV®, and Braun will actively induce, encourage, and abet this infringement with knowledge that such conduct is in contravention of Plaintiffs' rights under the '012 patent, in violation of 35 U.S.C. § 271(b).

65. Upon information and belief, Braun had actual and constructive knowledge of the application that later issued as the '012 patent prior to filing NDA No. 204957 and acted without a reasonable basis for a good faith belief that they would not be liable for infringing the '012 patent upon its issuance.

COUNT II
(DECLARATORY JUDGMENT OF INFRINGEMENT OF THE '012 PATENT)

66. Plaintiffs incorporate each of the preceding paragraphs 1 to 65 as if fully set forth herein.

67. This claim arises under the Declaratory Judgment Act, 28 U.S.C. §§ 2201 and 2202.

68. Plaintiffs are entitled to a declaration that, if Braun, prior to patent expiry, commercially manufactures, uses, offers for sale, or sells Braun's Generic Product within the United States, imports Braun's Generic Product into the United States, or induces such conduct, Braun would infringe the '012 patent under 35 U.S.C. § 271(a) and/or (b).

69. An actual controversy has arisen and now exists between the parties concerning whether Braun will directly or indirectly infringe the '012 patent.

70. Plaintiffs are entitled to an injunction restraining and enjoining Defendant and its officers, agents, attorneys and employees, and those acting in privity or concert with it, from engaging in the commercial manufacture, use, offer to sell, or sale within the United States, or importation into the United States of any of Braun's Generic Product until the expiration of the '012 patent, including any extensions and/or additional periods of exclusivity to which Plaintiffs are or become entitled.

71. Plaintiffs will be irreparably harmed by Braun's infringing activities unless those activities are enjoined by this Court. Plaintiffs do not have an adequate remedy at law.

COUNT III
(INFRINGEMENT OF THE '265 PATENT)

72. Plaintiffs incorporate each of the preceding paragraphs 1 to 71 as if fully set forth herein.

73. Braun's submission of NDA No. 204957 constitutes infringement of the '265 patent pursuant to 35 U.S.C. § 271(e)(2).

74. Upon information and belief, upon FDA approval of NDA No. 204957, Braun will induce infringement of at least Claim 7 of the '265 patent by making, using, offering to sell, or selling Braun's Generic Product in the United States, and/or importing Braun's Generic Product into the United States, in violation of 35 U.S.C. § 271.

75. Upon information and belief, upon FDA approval of NDA No. 204957, doctors, nurses, and other medical professionals will directly infringe at least Claim 7 of the '265 patent by using Braun's Generic Product, in violation of 35 U.S.C. § 271(a). Braun's Generic Product will be administered to treat pain in a human subject weighing at least 50 kg by co-administering to the subject a therapeutically effective amount of a first pharmaceutical composition comprising about 650 mg of acetaminophen and a therapeutically effective amount of a second pharmaceutical composition comprising an opioid analgesic, wherein the first pharmaceutical composition is administered to the subject intravenously, which co-administration will constitute direct infringement of at least Claim 7 of the '265 patent.

76. Upon information and belief, this direct infringement will occur at Braun's active behest, and with Braun's intent, knowledge, and encouragement. Braun will intentionally encourage infringement of at least Claim 7 of the '265 patent at least by way of the labeling for Braun's Generic Product which will contain recommendations and/or instructions for treating pain in a human subject weighing at least 50 kg by co-administering to the subject a

therapeutically effective amount of a first pharmaceutical composition comprising about 650 mg of acetaminophen and a therapeutically effective amount of a second pharmaceutical composition comprising an opioid analgesic, wherein the first pharmaceutical composition is administered to the subject intravenously.

77. Upon information and belief, Braun is aware of the '265 patent, which is listed in the Orange Book with respect to OFIRMEV®, and Braun will actively induce, encourage, and abet this infringement with knowledge that such conduct is in contravention of Plaintiffs' rights under the '265 patent, in violation of 35 U.S.C. § 271(b).

COUNT IV
(DECLARATORY JUDGMENT OF INFRINGEMENT OF THE '265 PATENT)

78. Plaintiffs incorporate each of the preceding paragraphs 1 to 77 as if fully set forth herein.

79. This claim arises under the Declaratory Judgment Act, 28 U.S.C. §§ 2201 and 2202.

80. Plaintiffs are entitled to a declaration that, if Braun, prior to patent expiry, commercially manufactures, uses, offers for sale, or sells Braun's Generic Product within the United States, imports Braun's Generic Product into the United States, or induces such conduct, Braun would infringe the '265 patent under 35 U.S.C. § 271(a) and/or (b).

81. An actual controversy has arisen and now exists between the parties concerning whether Braun will directly or indirectly infringe the '265 patent.

82. Plaintiffs are entitled to an injunction restraining and enjoining Defendant and its officers, agents, attorneys and employees, and those acting in privity or concert with it, from engaging in the commercial manufacture, use, offer to sell, or sale within the United States, or importation into the United States of any of Braun's Generic Product until the expiration of the

'265 patent, including any extensions and/or additional periods of exclusivity to which Plaintiffs are or become entitled.

83. Plaintiffs will be irreparably harmed by Braun's infringing activities unless those activities are enjoined by this Court. Plaintiffs do not have an adequate remedy at law.

PRAYER FOR RELIEF

WHEREFORE, Plaintiffs respectfully request the following relief:

- A. A judgment that Defendant infringed and is infringing each of the patents-in-suit;
- B. An order issued pursuant to 35 U.S.C. § 271(e)(4) that the effective date of any approval of Defendant's NDA No. 204957 shall not be earlier than the expiration date of the patents-in-suit, including any extensions and/or additional periods of exclusivity to which Plaintiffs are or become entitled;
- C. A declaration that if Defendant, prior to patent expiry, commercially manufactures, uses, offers for sale, or sells Braun's Generic Product within the United States, imports Braun's Generic Product into the United States, or induces such conduct, Defendant would infringe the patents-in-suit;
- D. A preliminary and permanent injunction restraining and enjoining Defendant and its officers, agents, attorneys and employees, and those acting in privity or concert with it, from engaging in the commercial manufacture, use, offer to sell, or sale within the United States, or importation into the United States of any of Braun's Generic Product until the expiration of the patents-in-suit, including any extensions and/or additional periods of exclusivity to which Plaintiffs are or become entitled;
- E. That Plaintiffs be awarded monetary relief if Defendant commercially manufactures, uses, offers for sale, or sells its generic version of Plaintiffs' OFIRMEV® brand product, or any other product that infringes or induces the infringement of the patents-in-suit, within the United States before the latest expiration date of the patents-in-suit, including any extensions and/or additional periods of exclusivity to which Plaintiffs are or becomes entitled;
- F. A declaration that this is an exceptional case and an award of attorneys' fees pursuant to 35 U.S.C. § 285;

- G. An award of costs and expenses in this action; and
- H. Such other and further relief as the Court may deem just and proper.

Dated: June 1, 2017



Keith R. Dutill (PA ID No. 46387)
Marissa R. Parker (PA ID No. 206162)
Chelsea A. Biemiller (PA ID No. 319625)
Stradley Ronon Stevens & Young, LLP
A Pennsylvania Limited Liability Partnership
2005 Market Street, Suite 2600
Philadelphia, PA 19103-7018
Tel: 215.564.8000
Fax: 215.564.8120
Attorneys for Mallinckrodt IP and Mallinckrodt Hospital Products Inc.

OF COUNSEL:

Kenneth G. Schuler
Marc N. Zubick
Sarah W. Wang
LATHAM & WATKINS LLP
330 North Wabash Avenue, Suite 2800
Chicago, IL 60611
(312) 876-7700

Daniel G. Brown
LATHAM & WATKINS LLP
885 Third Avenue
New York, NY 10022
(212) 906-1200

Gregory K. Sobolski
LATHAM & WATKINS LLP
505 Montgomery Street, Suite 2000
San Francisco, CA 94111
(415) 391-0600

Michelle R. Ma
LATHAM & WATKINS LLP
140 Scott Drive
Menlo Park, CA 94025
(650) 328-4600

Elissa N. Knoff

LATHAM & WATKINS LLP
555 Eleventh St, NW, Suite 1000
Washington, D.C. 20004
(202) 637-2200

*Attorneys for Mallinckrodt IP and
Mallinckrodt Hospital Products Inc.*

EXHIBIT A

(12) **United States Patent**
Royal et al.(10) **Patent No.:** **US 9,399,012 B2**
(45) **Date of Patent:** **Jul. 26, 2016**(54) **REDUCED DOSE INTRAVENOUS ACETAMINOPHEN**(75) Inventors: **Mike Allan Royal**, San Diego, CA (US); **James Bradley Breitmeyer**, Rancho Santa Fe, CA (US)(73) Assignee: **MALLINCKRODT IP**, Dublin (IE)

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Related U.S. Application Data

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(58) **Field of Classification Search**USPC 514/629
See application file for complete search history.(56) **References Cited**

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Primary Examiner — San-Ming Hui(74) *Attorney, Agent, or Firm* — Mayer Brown LLP(57) **ABSTRACT**

Described herein are compositions and methods for intravenous administration of acetaminophen at a single dose level of less than about 1000 mg for the treatment or prevention of pain (e.g., postoperative pain) and/or fever.

US 9,399,012 B2

Page 2

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US 9,399,012 B2

1

**REDUCED DOSE INTRAVENOUS
ACETAMINOPHEN****CROSS-REFERENCE TO RELATED
APPLICATIONS**

This application is related to and claims priority to U.S. Provisional Patent Application No. 60/987,761, entitled "Reduced Dose Intravenous Acetaminophen" filed on Nov. 13, 2007, which is incorporated herein by reference in its entirety.

BACKGROUND OF THE INVENTION

In the hospital, particularly in the postoperative setting, pain is a primary concern of patients. Opioid analgesics have been used to treat postoperative pain since 1784 and parenteral morphine has been a primary treatment modality since the 1850s. While opioids are highly effective in the treatment of many painful conditions, they have side effects and dose-dependent risks including nausea, vomiting, constipation, urinary retention, sedation, and respiratory depression. Similarly, non steroidal anti-inflammatory drugs (NSAIDs), including the older non selective (dual inhibitor) products and newer cyclo-oxygenase (COX)-2 products, have a variety of unwanted side effects especially when used in the perioperative setting. Non selective NSAIDs are associated with platelet dysfunction and the potential for bleeding at the surgical site, upper gastrointestinal ulcers and bleeding, edema, hypertension, congestive heart failure, renal dysfunction, severe skin reactions such as Stevens-Johnson syndrome and toxic epidermal necrolysis, anaphylaxis, and most recently, an increased risk of thrombotic cardiovascular events.

SUMMARY OF THE INVENTION

Described herein are pharmaceutical compositions having a reduced dose of acetaminophen for intravenous administration, and methods of using these compositions for treating and/or preventing pain and/or fever in a subject.

In some embodiments, the pharmaceutical compositions described herein comprise less than about 1 gram of acetaminophen, wherein the pharmaceutical composition is provided as a formulation suitable for intravenous administration. For example, various embodiments may comprise about 500 mgs to about 1 gram, or about 500 mgs to about 800 mgs, or about 500 mgs to about 750 mgs. In various embodiments, the pharmaceutical compositions described herein comprise about 600 mg to about 700 mg of acetaminophen.

In some embodiments, the pharmaceutical compositions described herein further comprise at least one antioxidant. In some embodiments, the at least one antioxidant comprises cysteine hydrochloride monohydrate, thioglycolic acid, thioglycolic acid, dithiothreitol, reduced glutathione, thiourea, alpha-thioglycerol, cysteine, acetyl cysteine, or mercaptoethane sulfonic acid, ascorbic acid ascorbic acid derivatives, an organic compound having at least one thiol, an alkyl polyhydroxylated compound, or a cycloalkyl polyhydroxylated compound.

In some embodiments, the pharmaceutical composition further comprises a buffering agent (e.g., disodium phosphate dehydrate). In some embodiments, the pharmaceutical composition has a pH from about 4 to about 8 when in solution. In some embodiments, the pharmaceutical composition has a pH of about 5 to about 6 when in solution.

2

In some embodiments, the pharmaceutical composition has an osmolality from about 250 mOsm/L to about 400 mOsm/L when in solution. In some embodiments, the pharmaceutical composition further comprises an isotonicity agent. In some embodiments, the isotonicity agent is dextrose, mannitol, or lactose.

In some embodiments, the pharmaceutical composition further comprises at least one analgesic agent other than acetaminophen. In some embodiments, the at least one analgesic agent other than acetaminophen comprises an anilide, an opioid, an NSAID, a cannabinoid, a pyrazolone, or a barbiturate.

In some embodiments, the pharmaceutical composition further comprises EDTA.

In a further aspect provided herein is a method for preventing or reducing pain or fever in a subject in need thereof, comprising administering to the subject, by an intravenous route of administration, a pharmaceutical compositions described herein. In some embodiments, the administration is repeated at least once with an interval of about 3 to about 5 hours. In some embodiments, the administration is repeated at least six times in a period of twenty four hours. In various embodiments, the administration is repeated three to eight times (e.g., 3 times, 4 times, five times, six times, seven times, or eight times) in a period of twenty four hours and about 3 to about 5 grams of acetaminophen (e.g., about 3 grams, about 4 grams or about 5 grams) is delivered over the twenty four hour period. In other embodiments, the administration is repeated three to eight times in a period of twenty four hours and less than about 4 grams of acetaminophen is delivered over the twenty four hour period.

In some embodiments, the pharmaceutical formulation for IV administration is a solution comprising: about 600 mg to about 700 mg of acetaminophen, cysteine hydrochloride monohydrate, disodium phosphate dehydrate, and mannitol, wherein the solution has a pH of between about 5 and about 6 and an osmolality of between about 200-400 mOsm/L. In some embodiments, the pharmaceutical composition in solution has an acetaminophen concentration of about 0.5% (w/v) to about 10% (w/v). In some embodiments, the acetaminophen concentration is about 1% (w/v). In some embodiments, the pharmaceutical composition to be administered further comprises EDTA.

In some embodiments, the subject to be treated is suffering from an infection. In some embodiments, the subject to be treated is suffering from a fever. In some embodiments, the subject to be treated is unconscious, sedated, fasting, nauseous, or unable to be administered a pharmaceutical composition by an oral route.

In some embodiments, the pharmaceutical composition is administered to the subject after a surgical intervention. In some embodiments, the pharmaceutical composition is administered within three hours of a surgical intervention on the subject. In some embodiments, the pharmaceutical composition is administered within 1 hour of a surgical intervention on the subject. In some embodiments, the pharmaceutical composition is administered postoperatively. In some embodiments, the subject to be treated is suffering from post-operative pain.

In various embodiments the pharmaceutical compositions described herein are administered as a pretreatment.

In another aspect provided herein is a method for preventing or reducing pain or fever in a subject in need thereof, comprising administering to the subject, by an intravenous route of administration, a pharmaceutical composition described herein.

US 9,399,012 B2

3

INCORPORATION BY REFERENCE

All publications and patent applications mentioned in this specification are herein incorporated by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference.

DETAILED DESCRIPTION OF THE INVENTION

To date, the standard intravenous (IV) dose of acetaminophen for the relief of pain or fever has been 1000 mg in adults and adolescents weighing at least 50 kg. At this dose level, the frequency of acetaminophen administration is limited to a maximum of once every six hours (i.e., four administrations per twenty four hours) to minimize the potential for hepatotoxicity. On the other hand, it has generally been noted that acetaminophen has greatest efficacy during its initial rise in plasma concentration, i.e., during the first few hours post-administration, and is less effective later on after the plasma concentration of the drug drops from its peak. While not wishing to be bound by theory, it is thought that this change in efficacy is likely due to a time and concentration-dependent modulation of the central and peripheral nociceptive pathways through which acetaminophen acts.

Further, if the duration of effect of a 1000 mg dose of acetaminophen is shorter in duration than 6 hours, the use of this dose is limited since dosing more frequently than every 6 hours, e.g., every 4 hours, leaves a gap in coverage due to the 4 g acetaminophen maximum daily limit. In the treatment of fever, a dose less than 1000 mg may be fully effective due to the fact that a lower plasma level (compared to that needed for pain) is needed to effectively reduce fever.

Thus, intravenous administration of a reduced dose of acetaminophen, as described herein, permits more frequent IV acetaminophen administration to yield better overall relief of symptoms for many patients while avoiding any potential gap artificially created by the daily limit.

Also, the reduced acetaminophen IV dose affords greater flexibility to the physician in customizing treatments to the needs of the patient, selecting the dose of other drugs for use in combination therapies and allowing for smoother transitions to oral products containing acetaminophen.

Accordingly, described herein are reduced IV dose formulations of acetaminophen for intravenous administration and the use of reduced IV doses of acetaminophen for use for the treatment or prevention of pain and/or fever.

Certain Terminology

Unless defined otherwise, all technical and scientific terms used herein have the same meaning as is commonly understood by one of skill in the art to which the claimed subject matter belongs. In the event that there is a plurality of definitions for terms herein, those in this section prevail. Where reference is made to a URL or other such identifier or address, it is understood that such identifiers can change and particular information on the internet can come and go, but equivalent information can be found by searching the internet or other appropriate reference source. Reference thereto evidences the availability and public dissemination of such information.

It is to be understood that the foregoing general description and the following detailed description are exemplary and explanatory only and are not restrictive of any subject matter claimed. In this application, the use of the singular includes the plural unless specifically stated otherwise. It must be noted that, as used in the specification and the appended claims, the singular forms "a", "an" and "the" include plural referents unless the context clearly dictates otherwise. It

4

should also be noted that use of "or" means "and/or" unless stated otherwise. Furthermore, use of the term "including" as well as other forms, such as "include", "includes", and "included" is not limiting.

5 Definition of standard chemistry terms may be found in reference works, including Carey and Sundberg "ADVANCED ORGANIC CHEMISTRY 4TH ED." Vols. A (2000) and B (2001), Plenum Press, New York. Unless otherwise indicated, conventional methods of mass spectroscopy, NMR, HPLC, IR and UV/Vis spectroscopy and pharmacology, within the skill of the art are employed. Unless specific definitions are provided, the nomenclature employed in connection with, and the laboratory procedures and techniques of, analytical chemistry, synthetic organic chemistry, and medicinal and pharmaceutical chemistry described herein are those known in the art. Standard techniques can be used for chemical syntheses, chemical analyses, pharmaceutical preparation, formulation, and delivery, and treatment of patients. Reactions and purification techniques can be performed e.g., using kits of manufacturer's specifications or as commonly accomplished in the art or as described herein. The foregoing techniques and procedures can be generally performed of conventional methods well known in the art and as described in various general and more specific references that are cited and discussed throughout the present specification. Throughout the specification, groups and substituents thereof can be chosen by one skilled in the field to provide stable moieties and compounds.

30 The terms "treat," "treating" or "treatment," and other grammatical equivalents as used herein, include alleviating, abating or ameliorating a disease or condition symptoms, preventing additional symptoms, ameliorating or preventing the underlying metabolic causes of symptoms, inhibiting the disease or condition, e.g., arresting the development of the disease or condition, relieving the disease or condition, causing regression of the disease or condition, relieving a condition caused by the disease or condition, or stopping the symptoms of the disease or condition, and are intended to include prophylaxis. The terms further include achieving a therapeutic benefit and/or a prophylactic benefit. By therapeutic benefit is meant eradication or amelioration of the underlying disorder being treated. Also, a therapeutic benefit is achieved with the eradication or amelioration of one or more of the physiological symptoms associated with the underlying disorder such that an improvement is observed in the patient, notwithstanding that the patient may still be afflicted with the underlying disorder. For prophylactic benefit, the compositions may be administered to a patient at risk of developing a particular disease, or to a patient reporting one or more.

35 The terms "effective amount," "therapeutically effective amount" or "pharmaceutically effective amount" as used herein, refer to a sufficient amount of at least one agent or compound being administered which will relieve to some extent one or more of the symptoms of the disease or condition being treated. The result can be reduction and/or alleviation of the signs, symptoms, or causes of a disease, or any other desired alteration of a biological system. For example, an "effective amount" for therapeutic uses is the amount of the composition comprising the compound as disclosed herein required to provide a clinically significant decrease in pain. An additional example is that an "effective amount" may be a dosage that decreases a fever. An appropriate "effective" amount in any individual case may be determined using techniques, such as a dose escalation study.

40 The terms "administer," "administering," "administration," and the like, as used herein, refer to the methods that may be used to enable delivery of compounds or compositions to the desired site of biological action. These methods

US 9,399,012 B2

5

include, but are not limited to oral routes, intraduodenal routes, parenteral injection (including intravenous, subcutaneous, intraperitoneal, intramuscular, intravascular or infusion), topical and rectal administration. Those of skill in the art are familiar with administration techniques that can be employed with the compounds and methods described herein, e.g., as discussed in Goodman and Gilman, *The Pharmacological Basis of Therapeutics*, current ed.; Pergamon; and Remington's, *Pharmaceutical Sciences* (current edition), Mack Publishing Co., Easton, Pa. In preferred embodiments, the compositions comprising acetaminophen as described herein are administered intravenously.

The term "acceptable" as used herein, with respect to a formulation, composition or ingredient, means having no persistent detrimental effect on the general health of the subject being treated.

The term "antioxidant" refers to a compound that prevents oxygen or oxygen-derived free radicals from interacting with other substances. Antioxidants are added to minimize or retard oxidative processes that occur with some drugs or excipients upon exposure to oxygen or in the presence of free radicals. These processes can often be catalyzed by light, temperature, hydrogen or concentration, presence of trace metals or peroxides.

The terms "enhance" or "enhancing," as used herein, means to increase or prolong either in potency or duration a desired effect. Thus, in regard to enhancing the effect of therapeutic agents, the term "enhancing" refers to the ability to increase or prolong, either in potency or duration, the effect of other therapeutic agents on a system. An "enhancing-effective amount," as used herein, refers to an amount adequate to enhance the effect of another therapeutic agent in a desired system.

"Concurrent administration," "administered in combination" or similar phrases referring to the acetaminophen and at least one additional component means that the components are administered concurrently to the mammal being treated. By "concurrently," it is meant that each component may be administered at the same time or sequentially in any order at different points in time. However, if not administered at the same time, they should be administered sufficiently closely in time so as to provide the desired enhancement of treatment effect. Suitable dosing intervals and the order of administration with such compounds will be readily apparent to those skilled in the art, once armed with the present disclosure. Preferably both components are administered at the same time or within the same hour.

As used herein, the term "animal" shall refer to a vertebrate animal. More preferably, the vertebrate animal is a mammal. As used herein, the term "mammal" shall refer to the *Mammalia* class of higher vertebrates. The term "mammal" includes, but is not limited to, a human.

As used herein, the term "pain" shall refer to all types of pain, including, but not limited, to nociceptive pain, neuropathic pain, post-operative pain, lower back pain, cluster headaches, herpes neuralgia, phantom limb pain, central pain, dental pain, opioid-resistant pain, visceral pain, surgical pain, bone injury pain, pain during labor and delivery, pain resulting from burns, including sunburn, post partum pain, migraine, and genitourinary tract-related pain including cystitis. Levels of pain in a subject can be quantified using standard subjective assay scales of pain including, e.g., the Pain Intensity Visual Analogue Scale or Pain Intensity Categorical Scale. Likewise, levels of "pain relief" can also be quantified by a subjective assay, e.g., Time to Perceptible and Meaningful Pain Relief.

6

The terms "intravenous formulation," or "intravenous acetaminophen formulation" shall refer to a single dose formulation of acetaminophen that is provided as a lyophilized powder (or other solid form) that, once reconstituted in solution, is physiologically compatible with intravenous administration (e.g., by injection, infusion or otherwise). Alternatively, the terms refer to a formulation that is provided as a solution.

Reduced Dose Acetaminophen Formulations for Intravenous Administration (IV Formulations)

In some embodiments, the IV acetaminophen formulations described herein are in the form of a lyophilized powder to be reconstituted in solution under sterile conditions prior to administration. In other embodiments, the IV acetaminophen formulations are provided as sterile solutions ready for administration. Appropriate containers (e.g., vials, bottles, ampules, containers, etc.) for the IV formulations in either of the forms just described, as well as aseptic techniques are well known.

20 IV Acetaminophen Dosage

In various embodiments, the single dose IV acetaminophen formulation contains less than about 1 gram of acetaminophen. In some embodiments, the single dose IV acetaminophen contains about 500 to about 1000 mgs. In some embodiments, the single dose IV acetaminophen contains about 550 mgs to about 900 mgs. In some embodiments, the single dose IV acetaminophen formulations described herein contain about 550 mg to about 800 mg of acetaminophen, i.e., about 560 mg, 570 mg, 580 mg, 600 mg, 610 mg, 620 mg, 630 mg, 640 mg, 650 mg, 660 mg, 670 mg, 675 mg, 680 mg, 690 mg, 700 mg, 720 mg, 750 mg, 775 mg, or any other amount of acetaminophen from about 550 mg to about 800 mg of acetaminophen. In some embodiments, an IV acetaminophen formulation contains about 600 mg to about 700 mg of acetaminophen, i.e., about 610 mg, 620 mg, 630 mg, 640 mg, 650 mg, 660 mg, 670 mg, 680 mg, 690 mg, or any other amount of acetaminophen from about 600 mg to about 700 mg of acetaminophen. In one embodiment, the acetaminophen formulation contains about 650 mg of acetaminophen.

In some embodiments, the concentration of acetaminophen in an IV formulation solution described herein is about 0.3% (w/v) to about 12% (w/v), i.e., about 0.4%, 0.5%, 0.6%, 0.7%, 0.8%, 0.9%, 1.0%, 1.1%, 1.2%, 1.5%, 2.5%, 3%, 3.7%, 4%, 4.5%, 5%, 6%, 7%, 8%, 8.5%, 9%, 10%, 10.5%, 11%, or any other concentration from about 0.3% (w/v) to about 12% (w/v). In some embodiments the concentration of acetaminophen is about 0.7% (w/v) to about 1.4% (w/v), i.e., about 0.8%, 0.9%, 1.0%, 1.1%, 1.2%, 1.3% or any other concentration of acetaminophen from about 0.7% (w/v) to about 1.4% (w/v). In one embodiment, the concentration of acetaminophen is about 1.0% (w/v).

In some embodiments, the volume of an IV acetaminophen formulation solution is about 30 to about 200 ml, i.e., about 30, 35, 40, 45, 55, 60, 65, 75, 80, 85, 90, 92, 95, 100, 105, 110, 125, 130, 150, 175, 180, or another volume of IV formulation solution from about 30 to about 200 ml. In some embodiments, the volume of the IV formulation is about 75 to about 125 ml. In another embodiment the volume is about 40 to about 75 ml. In one embodiment, the volume of the IV formulation is about 100 ml.

Antioxidants

Generally, the acetaminophen formulations described herein also contains at least one antioxidant to increase the stability of acetaminophen in solution. Examples of suitable antioxidants include, but are not limited to, cysteine hydrochloride monohydrate, thioglycolic acid, thiolacetic acid,

US 9,399,012 B2

7

dithiothreitol, reduced glutathione, thiourea, alpha-thioglycerol, cysteine, acetylcysteine, methionine, mercaptoethane sulfonic acid, metabisulfite, ascorbic acid ascorbic acid derivatives (e.g., ascorbyl palmitate), sodium citrate, an organic compound having at least one thiol, an alkyl polyhydroxylated compound, a cycloalkyl polyhydroxylated compound, a hydroxypolycarboxylic acid, an alpha-hydroxypolycarboxylic acid (e.g., citric acid), tocotrienol, dimethyl glycine, betaine, butylated hydroxyanisole, butylated hydroxytoluene, tocopherol, tocopherol polyethylene glycol succinate, butylated hydroxytoluene (BHT), butylated hydroxyanisole (BHA), propyl gallate, hydroquinone, hydroxycoumarins, ethanolamine, lecithin, cephalin, malic acid, sorbitol, phosphoric acid, thiadipropionic acid and its esters, dithiocarbamates or any combination thereof. In one embodiment, the acetaminophen formulation is free of polyethylene glycol or a derivative thereof. In another embodiment, the acetaminophen formulation is free of sulfites. In one embodiment, the antioxidant is cysteine hydrochloride monohydrate. In yet another embodiment, the antioxidant is mannitol.

In some embodiments, the amount % (w/w) of the antioxidant in the solid form of the IV formulation (i.e., prior to preparation in solution) is about 0.10% (w/w) to about 5.0% (w/w), i.e., 0.15% (w/w), 0.17% (w/w), 0.20% (w/w), 0.30% (w/w), 0.40% (w/w), 0.45% (w/w), 0.50% (w/w), 0.52% (w/w), 0.55% (w/w), 0.60% (w/w), 0.70% (w/w), 0.80% (w/w), 1.0% (w/w), 1.3% (w/w), 1.5% (w/w), 1.7% (w/w), 2.0% (w/w), 2.2% (w/w), 2.3% (w/w), 2.5% (w/w), 2.7%, 2.8%, 3.0% (w/w), 3.2%, 3.5% (w/w), 3.6% (w/w), 4.0% (w/w), 4.7% (w/w), or any other amount of antioxidant % (w/w) from about 0.10% (w/w) to about 5.0% (w/w). In some embodiments, the amount % (w/w) of antioxidant is about 0.30% (w/w) to about 1.0% (w/w). In one embodiment, the amount % (w/w) of antioxidant is about 0.50% (w/w).

In some embodiments, the concentration of the antioxidant in an IV formulation solution prior to administration ranges from about 0.01 mg/ml to about 10 mg/ml, i.e., 0.02 mg/ml, 0.03 mg/ml, 0.05 mg/ml, 0.08 mg/ml, 0.09 mg/ml, 0.10 mg/ml, 0.12 mg/ml, 0.13 mg/ml, 0.15 mg/ml, 0.18 mg/ml, 0.20 mg/ml, 0.22 mg/ml, 0.25 mg/ml, 0.27 mg/ml, 0.30 mg/ml, 0.40 mg/ml, 0.45 mg/ml, 0.50 mg/ml, 0.60 mg/ml, 0.80 mg/ml, 1.2 mg/ml, 1.5 mg/ml, 2.0 mg/ml, 2.5 mg/ml, 3.0 mg/ml, 3.5 mg/ml, 4.0 mg/ml, 5.0 mg/ml, 6.0 mg/ml, 7.5 mg/ml, 8.0 mg/ml, 9 mg/ml, 9.5 mg/ml, or any other concentration of antioxidant from about 0.01 mg/ml to about 10 mg/ml. In some embodiments, the concentration of antioxidant is about 0.08 mg/ml to about 0.50 mg/ml. In one embodiment, the concentration of antioxidant is about 0.25 mg/ml.

Buffering Agents

In some embodiments, an IV acetaminophen formulation contains at least one buffering agent to maintain the pH of the formulation within an acceptable range as described herein. The buffer used is a buffer compatible with parenteral administration in humans, the pH of which may be adjusted between 4 and 8. In some embodiments, the pH of an IV acetaminophen formulation is from about pH 4 to about pH 8, i.e., pH 4.5, pH 4.6, pH 4.8, pH 5.0, pH 5.5, pH 6.2, pH 6.5, pH 7.5, or any other pH value from about pH 4 to about pH 8. In some embodiments, the pH of the IV acetaminophen formulation is from about pH 5 to about pH 7.0, i.e., about pH 5.2, pH 5.5, pH 5.6, pH 6.0, pH 6.4, or any other pH value from about pH 5 to about pH 7.0. In one embodiment, the IV acetaminophen formulation has a pH of about 5 to about 6.

In some embodiments, buffering agents have a pKa from about 4.5 to about 6.5, i.e., 4.6, 4.8, 5.0, 5.2, 5.3, 5.4, 5.5, 5.8, 6.0, 6.2, 6.4, or any other pKa from about 4.5 to about 6.5.

8

In some embodiments, the buffering agent is a pharmaceutically acceptable salt or acid of citrate, phosphate, acetate, glutamate, tartrate, benzoate, lactate, histidine or other amino acids, gluconate, malate, succinate, formate, propionate, carbonate, or any combination thereof adjusted to an appropriate pH, as described herein, with acid (e.g., hydrochloric acid) or base (e.g., sodium hydroxide) as required. In one embodiment, the buffering agent is disodium phosphate dehydrate.

In some embodiments, the amount % (w/w) of the buffering agent in the solid form of the IV formulation (i.e., prior to preparation in solution) is about 0.05% (w/w) to about 2% (w/w), i.e., about 0.08% (w/w), 0.10% (w/w), 0.15% (w/w), 1.0% (w/w), 1.3% (w/w), 1.5% (w/w), 1.7% (w/w), 0.20% (w/w), 0.22% (w/w), 0.25% (w/w), 0.26% (w/w), 0.27% (w/w), 0.28% (w/w), 0.30% (w/w), 0.35% (w/w), 0.40% (w/w), 0.50% (w/w), 0.60% (w/w), 0.70% (w/w), 0.80% (w/w), 1.2% (w/w), 1.4% (w/w), 1.5% (w/w), 1.7%, or any other amount of buffering agent % (w/w) from about 0.05% (w/w) to about 2.0% (w/w). In some embodiments, the amount % (w/w) of the buffering agent is about 0.10% to about 0.70%. In one embodiment, the amount % (w/w) of the buffering agent is about 0.26%.

In some embodiments, the concentration of the buffering agent in an IV formulation solution prior to administration ranges from about 0.01 mg/ml to about 10 mg/ml, i.e., 0.02 mg/ml, 0.03 mg/ml, 0.05 mg/ml, 0.08 mg/ml, 0.09 mg/ml, 0.10 mg/ml, 0.12 mg/ml, 0.13 mg/ml, 0.15 mg/ml, 0.17 mg/ml, 0.20 mg/ml, 0.22 mg/ml, 0.25 mg/ml, 0.27 mg/ml, 0.30 mg/ml, 0.35 mg/ml, 0.40 mg/ml, 0.45 mg/ml, 0.50 mg/ml, 0.60 mg/ml, 0.70 mg/ml, 0.80 mg/ml, 0.90 mg/ml, 1.0 mg/ml, 1.2 mg/ml, 1.5 mg/ml, 1.8 mg/ml, 2.0 mg/ml, 2.5 mg/ml, 3.0 mg/ml, 3.5 mg/ml, 4.0 mg/ml, 4.5 mg/ml, 5.0 mg/ml, 6.0 mg/ml, 7.5 mg/ml, 8.0 mg/ml, 9 mg/ml, 9.5 mg/ml, or any other concentration of buffering agent from about 0.01 mg/ml to about 10 mg/ml. In some embodiments, the concentration of buffering agent is about 0.08 mg/ml to about 0.30 mg/ml. In one embodiment, the concentration of buffering agent is about 0.13 mg/ml.

Isotonicity Agents

In some embodiments, an IV acetaminophen formulation also contains one or more isotonicity agents to maintain the osmolality of the formulation in a range that is physiologically compatible with IV administration. In some embodiments, the osmolality of the IV acetaminophen formulation is about 230 mOsm/L to about 420 mOsm/L, i.e., about 240 mOsm/L, 250 mOsm/L, 260 mOsm/L, 270 mOsm/L, 280 mOsm/L, 290 mOsm/L, 300 mOsm/L, 305 mOsm/L, 310 mOsm/L, 320 mOsm/L, 350 mOsm/L, 375 mOsm/L, 400 mOsm/L or any other osmolality from about 240 mOsm/L to about 420 mOsm/L. In some embodiments, the osmolality of the IV acetaminophen formulation is about 280 mOsm/L to about 320 mOsm/L, i.e., about 290 mOsm/L, 295 mOsm/L, 300 mOsm/L, 305 mOsm/L, 310 mOsm/L, 315 mOsm/L, or any other osmolality from about 280 mOsm/L to about 320 mOsm/L. In one embodiment, the osmolality of the IV acetaminophen formulation is about 200-400 mOsm/L.

Suitable agents for adjusting the isotonicity of IV acetaminophen formulations include, but are not limited to, mannitol, sorbitol, glycerol, sucrose, glucose, dextrose, levulose, fructose, lactose, polyethylene glycols 400 to 4000, phosphates, sodium chloride, potassium chloride, calcium chloride, calcium gluconoglucoheptonate, dimethyl sulfone. In one embodiment, the isotonicity agent is mannitol.

In some embodiments, the amount % (w/w) of the isotonicity agent in the solid form of the IV formulation (i.e., prior to preparation in solution) is about 5% (w/w) to about 95% (w/w), i.e., about 10% (w/w), 15% (w/w), 20% (w/w), 25% (w/w), 30% (w/w), 35% (w/w), 40% (w/w), 45% (w/w), 50% (w/w), 55% (w/w), 60% (w/w), 65% (w/w), 70% (w/w), 72% (w/w), 74% (w/w), 76% (w/w), 78% (w/w), 79% (w/w), 80%

US 9,399,012 B2

9

(w/w), 81% (w/w), 82% (w/w), 84% (w/w), 86% (w/w), 90% (w/w), 92% (w/w), or any other amount of isotonicity agent % (w/w) from about 5% (w/w) to about 95% (w/w). In some embodiments, the amount of isotonicity agent % (w/w) is about 65% (w/w) to about 85% (w/w). In one embodiment, the amount of isotonicity agent % (w/w) is about 79%.

In some embodiments, the concentration of the isotonicity agent in an IV formulation solution prior to administration ranges from about 1.0 mg/ml to about 150 mg/ml, i.e., 1.0 mg/ml, 2.0 mg/ml, 3.0 mg/ml, 3.5 mg/ml, 4.0 mg/ml, 4.5 mg/ml, 5.0 mg/ml, 8.0 mg/ml, 12 mg/ml, 15 mg/ml, 20 mg/ml, 25 mg/ml, 30 mg/ml, 32 mg/ml, 35 mg/ml, 37 mg/ml, 38 mg/ml, 40 mg/ml, 50 mg/ml, 60 mg/ml, 75 mg/ml, 80 mg/ml, 90 mg/ml, 95 mg/ml, 100, 110, 120, 140, or any other concentration of buffering agent from about 5 mg/ml to about 150 mg/ml. In some embodiments, the concentration of buffering agent is about 0.08 mg/ml to about 0.30 mg/ml. In one embodiment, the concentration of buffering agent is about 0.13 mg/ml.

Stabilizers

In some embodiments, IV acetaminophen formulations described herein also include a stabilizer, e.g., a chelating agent such as ethylene diamino tetraacetic acid (EDTA), ethylene diamino, N,N'-diacetic-N,N'-dipropionic acid, ethylene diamino tetraphosphonic acid, 2,2'-(ethylene diamino) dibutyric acid, nitrilotriacetic acid, or ethylene-glycol bis (diaminoethyl ether) N,N,N',N'-tetraacetic acid and sodium or calcium salts thereof. In some embodiments, the IV acetaminophen formulation includes EDTA as the stabilizer.

In some embodiments, the IV acetaminophen formulations described herein contain a stabilizer in the amount of about 0.005 to about 1.0 mg/ml. In some embodiments, the stabilizer is present in an amount of about 0.01 mg/ml, 0.05 mg/ml, 0.1 mg/ml, 0.5 mg/ml, or 1.0 mg/ml.

In some embodiments, to reduce oxidation of acetaminophen in solution and thereby increase its stability, oxygen is removed from an IV formulation solution by bubbling an inert gas (e.g., argon or nitrogen) through the solution under sterile conditions. Methods for minimizing oxidative degradation of acetaminophen solutions during storage are described in further detail in, e.g., U.S. Pat. No. 6,992,218, which is incorporated herein by reference in its entirety.

Methods of Treatment

In many cases, IV administration of acetaminophen is considered the most suitable route of administration for expedient and efficacious relief of a patient's pain or fever, particularly in a hospital setting. In some embodiments, a subject to be administered an IV formulation of acetaminophen (e.g., an adult subject or adolescent weighing at least about 50 kg), as described herein, is unconscious, sedated, fasting, nauseous, or unable to be administered a pharmaceutical composition by an oral route. Additionally, the rectal route is associated with highly variable bioavailability and slow absorption, and in children, the efficacious rectal dose exposes some pediatric patients to a potentially toxic exposure. In some embodiments, a patient suffering from pain or fever is in need of a faster onset of pain relief or fever treatment than possible by acetaminophen administration through an administration route other than by an IV administration.

In some embodiments, the IV formulations described herein are used as a pretreatment to another therapy. In some of these embodiments, pretreatment with an IV formulation described herein allows the use of a lower dose of acetaminophen. In some embodiments, the IV formulation described herein is administered before chemotherapy treatment, radiation treatment, a biopsy, or a blood transfusion. It should be understood that these are non-limiting examples and that the

10

IV formulations described herein can be administered as a pretreatment to any therapy where pain and/or fever are predicted to occur.

The IV formulations described herein can be used for reducing pain conditions including, but not limited to, acute nociceptive pain, acute neuropathic pain, postoperative pain, lower back pain, cluster headaches, herpes neuralgia, phantom limb pain, central pain, dental pain, opioid-resistant pain, visceral pain, surgical pain, procedural pain, bone injury pain, 10 pain during labor and delivery, pain resulting from burns, post partum pain, headache, muscular aches, backache, arthritis pain, the common cold, toothache, dental pain, osteoarthritis pain, menstrual pain, menstrual cramps, migraine, and genitourinary tract-related pain including cystitis. In some 15 embodiments, the IV formulation is administered preemptively to a subject, i.e., prior to the onset of pain or a pain-inducing condition or stimulus (e.g., a surgical operation). In some embodiments, the IV formulations described herein are used to reduce fever, including, but not limited to, fever due to 20 infections, drug reactions, allergic reactions, transfusion reactions, stroke, surgery, heat stroke, rheumatic diseases, cancer, or fever of unknown origin. In some embodiments, the IV formulations described herein are administered to a patient undergoing a dental procedure.

25 In some embodiments, the IV formulation is administered to a subject after undergoing a surgical intervention, e.g., within about 12 hours after a surgical intervention, i.e., within 11 hours, 10 hours, 9 hours, 8 hours, 6 hours, 5 hours, 4 hours, 3 hours, 2 hours, 1 hours, 45 minutes, 30 minutes 15 minutes, 30 5 minutes, or any period within about 12 hours following a surgical intervention.

In some embodiments, a subject is administered the IV formulation prior to a surgical intervention, e.g., about 4 hours or less prior to the surgical intervention, i.e., about 3 hours, 2 hours, 1 hours, 30 minutes, 15 minutes or even during the surgical intervention itself.

35 Depending on the concentration of acetaminophen in an IV formulation solution and consistent with the acetaminophen dose levels described herein, the volume of IV formulation 40 solution to be administered can vary from about 1 mL to about 200 mL, e.g., 5 mL, 10 mL, 20 mL, 25 mL, 30 mL, 40 mL, 50 mL, 60 mL, 65 mL, 70 mL, 85 mL, 90 mL, 100 mL, 110 mL, 120 mL, 130 mL, 140 mL, 150 mL, 160 mL, 180 mL, or any other volume of IV formulation solution from about 1 mL to 45 about 200 mL.

In some embodiments, the amount of time required for 50 administration of the IV formulation ranges from about 1 minute to about 1 hours, i.e., about 5 minutes, 10 minutes, 11 minutes, 15 minutes, 20 minutes, 30 minutes, 45 minutes, or any other administration time from about 1 minute to about 1 hour. In some embodiments, the amount of time required for administration of the IV formulation ranges from about 5 minutes to about 45 minutes, or about 5 minutes to about 30 minutes, or about 5 minutes to about 15 minutes.

55 Depending on the severity and persistence of a subject's condition, and in accordance with a medical caregiver's judgment, an IV formulation dose of acetaminophen, as described herein, can be administered in an interval to allow for the administration of about 3 to about 5 grams in a 24 hour period.

60 In some embodiments, the IV formulation of acetaminophen is administered in an interval sufficient to allow for the administration of about 4 grams in a 24 hour period. In some embodiments, the IV formulation is administered between 1 to 6 times, i.e., 1, 2, 3, 4, 5, 6 times every twenty four hours, 65 as deemed necessary by a medical caregiver. In some embodiments, the frequency of administration is not greater than once every four hours.

11

In various embodiments, the IV formulation of acetaminophen is dosed so as to provide less than about 4 grams over a 24 hour period. In various embodiments, the IV formulation of acetaminophen is dosed three to six times in a 24 hour period. For example, in some embodiments, the IV formulation of acetaminophen is dosed three times in a 24 hour period. In other embodiments, the IV formulation of acetaminophen is dosed four times in a 24 hour period. In still other embodiments, the IV formulation of acetaminophen is dosed five times in a 24 hour period. In some embodiments, the IV formulation of acetaminophen is dosed six times in a 24 hour period. In some embodiments, the IV formulation of acetaminophen is dosed seven times in a 24 hour period. In some embodiments, the IV formulation of acetaminophen is dosed eight times in a 24 hour period.

Combination Therapies

The acetaminophen IV formulations described herein can also be used in combination with other therapeutic reagents, e.g., other analgesics, antipyretics, or anti-inflammatory agents that are selected for their therapeutic or palliative value. In general, where a combination therapy is employed, other agents do not have to be administered in the same pharmaceutical composition as acetaminophen, and may, because of different physical and chemical characteristics, be administered by different routes. The determination of the mode of administration and the advisability of administration, where possible, in the same pharmaceutical composition, is well within the knowledge of the skilled clinician with the teachings described herein. The initial administration of either the IV acetaminophen formulation or the one or more therapeutic agents (e.g., analgesic agents other than acetaminophen) to be used in combination with acetaminophen can be made according to established protocols known in the art, and then, based upon the observed effects, the dosage, modes of administration and times of administration can be modified by the skilled clinician.

The particular choice of compounds (e.g., analgesic agents) for use in combination with the IV acetaminophen formulation described herein will depend on the diagnosis of the attending physicians (or other medical caregivers) and their judgment of the condition of the patient and the appropriate treatment protocol. The compounds may be administered concurrently (e.g., simultaneously, essentially simultaneously or within the same treatment protocol) or sequentially, depending upon the severity of pain experienced by the patient, the nature of the disease, disorder, or condition, the condition, and the actual choice of compounds used. The determination of the order of administration, and the number of repetitions of administration of each therapeutic agent during a treatment protocol, is well within the knowledge of the skilled physician after evaluation of the disease being treated and the condition of the patient.

For combination therapies described herein, dosages of the compounds to be co-administered with an acetaminophen IV formulation will vary depending on the type of co-drug employed, on the amount of pain experienced by the patient, the risk for addiction, the disease or condition being treated and so forth. In addition, when co-administered with one or more biologically active agents, the acetaminophen IV formulation provided herein may be administered either simultaneously with the biologically active agent(s), or sequentially. If administered sequentially, the attending physician will decide on the appropriate sequence of administering protein in combination with the biologically active agent(s).

In any case, the multiple therapeutic agents (one of which is an acetaminophen IV formulation described herein) may be administered in any order or even simultaneously. If simulta-

12

neously, the multiple therapeutic agents may be provided in a single, unified IV form, or in multiple forms (by way of example only, either as a single IV formulation, as multiple IV formulations, or as IV formulation and a pill). One of the therapeutic agents may be given in multiple doses, or both may be given as multiple doses. If not simultaneous, the timing between the multiple doses may vary from more than 1 minute to less than 12 hours. In some embodiments, the timing between the multiple doses is from between about 1 minute to about 6 hours, or about 1 minute and about 3 hours, or about 1 minute and about 1 hour. In addition, the combination methods, compositions and formulations are not to be limited to the use of only two agents; the use of multiple therapeutic combinations is also envisioned.

The pharmaceutical agents which make up the combination therapy disclosed herein may be a combined dosage form (i.e., a combined IV formulation) or in separate dosage forms intended for substantially simultaneous administration. The pharmaceutical agents that make up the combination therapy may also be administered sequentially, with either therapeutic compound being administered by a regimen calling for two-step administration. The two-step administration regimen may call for sequential administration of the active agents or spaced-apart administration of the separate active agents. The time period between the multiple administration steps may range from, a few minutes to several hours, depending upon the properties of each pharmaceutical agent, such as potency, solubility, bioavailability, plasma half-life and kinetic profile of the pharmaceutical agent.

The compounds described herein and combination therapies can be administered before, during or after the occurrence of a fever or painful condition, and the timing of administering the composition containing a compound can vary. Thus, for example, the compounds can be used as a prophylactic and can be administered continuously to subjects with a propensity to develop conditions (e.g., body aches and chills following chemotherapy treatment) or diseases in order to prevent the occurrence of the disease or condition. The compounds and compositions can be administered to a subject during or as soon as possible after the onset of the symptoms. The administration of the compounds can be initiated within the first 48 hours of the onset of the symptoms, preferably within the first 48 hours of the onset of the symptoms, more preferably within the first 6 hours of the onset of the symptoms, and most preferably within 3 hours of the onset of the symptoms.

A compound is preferably administered as soon as is practicable before or after the onset of a painful condition (e.g., postoperative pain), and for a length of time necessary for the treatment of the disease, such as, for example, from about 1 month to about 3 months.

Exemplary Analgesic Agents for Use in Combination with an Acetaminophen IV Formulation**Opioids**

In some embodiments, an acetaminophen IV formulation described herein is used in any combination with one or more opioids, which include, but are not limited to allylprodine, alphamethylfentanyl, alfentanil, bezitramide, buprenorphine, butorphanol, carfentanyl, codeine, dextropropoxyphene, dextromoramide, dezocine, diacetylmorphine, dihydrocodeine, dipipanone, morphine, dihydrocodeine, etorphine, fentanyl, hydrocodone, hydromorphone, ketobemidone, lefetamine, levorphanol, levo-alphaacetylmethadol, levomethorphan, meptazinol, methadone, morphine, nalbuphine, nico-morphine, ohmesfentanyl, opium, oripavine, oxycodone, oxymorphone, methadone, PEPAP, pentazocine, pethidine,

US 9,399,012 B2

13

phenazocine, piritamide, prodine, propoxyphene napsylate, remifentanil, sufentanil, tilidine, thebaine, tramadol, and tapentadol.

Non-Steroidal Anti-Inflammatory Drugs (NSAIDs)

In some embodiments, an acetaminophen IV formulation described herein is used in any combination with one or more NSAIDs, which include, but are not limited to amoxiprin, benorilate, choline magnesium salicylate, diflusinal, fasilamine, methyl salicylate, magnesium salicylate, salicyl salicylate, diclofenac, aceclofenac, acetaminophen, bromfenac, ethenzamide, etodolac, indometacin, nabumetone, sulindac, tolmetin, carprofen, fenbufen, fenoprofen, flurbiprofen, ketoprofen, ketorolac, loxoprofen, naproxen, oxaprozin, tiaprofenic acid, suprofen, mefenamic acid, meclofenamic acid, phenylbutazone, metamizole, oxyphenbutazone, sulfipyrazone, piroxicam, lornoxicam, meloxicam, tenoxicam, nimesulide salixylates, arylalkanoic acids, 2-arylpropionic acids (profens), n-arylanthranilic acids (fenamic acids), pyrazolidine derivatives, oxicams, and COX-2 inhibitors.

Other Analgesic Agents

In some embodiments, an acetaminophen IV formulation described herein is used in any combination with one or more analgesic agents not described above, including, but not limited to, barbiturates (e.g., butalbital), pyrazolones (e.g., amiphenazone, metamizole, phenazone), cannabinoids (e.g., tetrahydrocannabinol), ziconotide, choline magnesium fentanyl, oxycodone, codeine, dihydrocodeine, dihydrocodeine enol acetate, morphine, desomorphine, apomorphine, diamorphine, pethidine, methadone, dextropropoxyphene, pentazocine, dextromoramide, oxymorphone, hydromorphone, dihydromorphone, noscapine, papaverine, papaveretum, alfentanil, buprenorphine, tramadol and pharmaceutically acceptable salts, derivatives, homologs or analogs thereof as well as opioid agonists.

Exemplary Antiemetic Agents for Use in Combination with an Acetaminophen IV Formulation

In some embodiments, an acetaminophen IV formulation described herein is used in any combination with one or more antiemetic agents not described above, including, but not limited to, antihistamines (e.g., Cyclizine, Diphenhydramine, Dimenhydrinate, Meclizine, Promethazine, Pentazine, Phenergan, Promacot, or Hydroxyzine); 5-HT₃ receptor antagonists (e.g., Dolasetron, Granisetron, Ondansetron, Tropisetron, or Palonosetron); and dopamine antagonists (e.g., Domperidone, Droperidol, haloperidol, chlorpromazine, promethazine, prochlorperazine, or metoclopramide).

Kits

In some embodiments provided herein are kits that can simplify the administration of an IV acetaminophen formulation to a patient. In some embodiments, a kit comprises a unit dosage form of an acetaminophen IV formulation as described herein provided as a sterile lyophilate to be reconstituted by addition of sterile water. In other embodiments, the

14

IV formulation is provided as a sterile degassed solution ready for administration. The kit can further comprise a label or printed instructions on the use of the acetaminophen IV formulation to treat pain or fever. The kit can also further comprise a unit dosage form of another therapeutic agent, for example, a container containing an effective amount of a second analgesic agent for use in combination with the acetaminophen IV formulation. In some embodiments, a kit further comprises a device that is useful for administering the IV formulation unit dosage forms. Examples of such a device include, but are not limited to, a syringe or a drip bag.

While preferred embodiments of the present invention have been shown and described herein, it will be understood that such embodiments are provided by way of example only. Numerous variations, changes, and substitutions can be made without departing from the scope of the invention. It should be understood that various alternatives to the embodiments of the invention described herein may be employed in practicing the invention. It is intended that the following claims define the scope of the invention and that methods and structures within the scope of these claims and their equivalents be covered thereby. Thus, these examples should not be read as limiting the example in any way. For example, different amounts of the components described in the following examples as well as the components themselves can be changed according to the disclosure provided herein.

EXAMPLES

Example 1

IV Acetaminophen Formulations

TABLE 1

Exemplary IV Formulation of Acetaminophen	
Acetaminophen	0.550 g-1.000 g
Excipients:	
Antioxidant	0.0100-0.0200 g
pH modulator(s)	qs pH 5-6
Buffer	0.005-0.01 g
Isotonic Agent	1.5-3.5 g
Solvent	qs 50.0-100.0 mL

Example 1A

IV Acetaminophen Formulations

Example 1A is prepared according to the procedure outlined in Example 2 using the amounts of the excipients described in Example 1.

Formula 1(A)					
Acetaminophen					
Excipients:	0.550 g	0.650 g	0.750 g	0.850 g	0.950 g
Antioxidant	Reduced Glutathione				
pH Modulator	Sodium hydroxide				
pH Modulator	Hydrochloric Acid				
Buffering Agent	Sodium Citrate				

US 9,399,012 B2

15

-continued

16

Formula 1(A)					
Acetaminophen					
Excipients:	0.550 g	0.650 g	0.750 g	0.850 g	0.950 g
Isotonicity Agent	Sodium Chloride				
Solvent	Sterile Water for injection				

Example 1B

IV Acetaminophen Formulations

15

Example 1B is prepared according to the procedure outlined in Example 2 using the amounts of the excipients described in Example 1.

Formula 1(B)					
Acetaminophen					
Excipients:	0.550 g	0.650 g	0.750 g	0.850 g	0.950 g
Antioxidant	Methionine	Methionine	Methionine	Methionine	Methionine
pH Modulator	Sodium hydroxide				
pH Modulator	Hydrochloric Acid				
Buffering Agent	Sodium Acetate				
Isotonicity Agent	Mannitol	Mannitol	Mannitol	Mannitol	Mannitol
Solvent	Sterile Water for injection				

Example 1C

IV Acetaminophen Formulations

40

Example 1C is prepared according to the procedure outlined in Example 2 using the amounts of the excipients described in Example 1.

Formula 1(C)					
Acetaminophen					
Excipients:	0.550 g	0.650 g	0.750 g	0.850 g	0.950 g
Antioxidant	Cysteine Hydrochloride	Cysteine Hydrochloride	Cysteine Hydrochloride	Cysteine Hydrochloride	Cysteine Hydrochloride
pH Modulator	Monohydrate Sodium hydroxide	Monohydrate hydroxide	Monohydrate hydroxide	Monohydrate hydroxide	Monohydrate hydroxide
pH Modulator	Hydrochloric Acid	Hydrochloric Acid	Hydrochloric Acid	Hydrochloric Acid	Hydrochloric Acid
Buffering Agent	Disodium Phosphate	Disodium Phosphate	Disodium Phosphate	Disodium Phosphate	Disodium Phosphate
Isotonicity Agent	Dehydrate Mannitol	Dehydrate Mannitol	Dehydrate Mannitol	Dehydrate Mannitol	Dehydrate Mannitol
Solvent	Sterile Water for injection	Sterile Water for injection	Sterile Water for injection	Sterile Water for injection	Sterile Water for injection

US 9,399,012 B2

17
Example 1D

18

IV Acetaminophen Formulations

Example 1D is prepared according to the procedure outlined in Example 2 using the amounts of the excipients described in Example 1. 5

Excipients:	Formula 1(D)				
	Acetaminophen				
	0.550 g	0.650 g	0.750 g	0.850 g	0.950 g
Antioxidant	Ascorbic Acid				
pH Modulator	Sodium hydroxide				
pH Modulator	Hydrochloric Acid				
Buffering Agent	Sodium Tartate				
Isotonicity Agent	Glycerol	Glycerol	Glycerol	Glycerol	Glycerol
Solvent	Sterile Water for injection				

Example 1E

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IV Acetaminophen Formulations

Example 1E is prepared according to the procedure outlined in Example 2 using the amounts of the excipients described in Example 1.

Excipients:	Formula 1(E)				
	Acetaminophen				
	0.550 g	0.650 g	0.750 g	0.850 g	0.950 g
Antioxidant	Acetylcysteine	Acetylcysteine	Acetylcysteine	Acetylcysteine	Acetylcysteine
pH Modulator	Sodium hydroxide				
pH Modulator	Hydrochloric Acid				
Buffering Agent	Disodium Phosphate				
Isotonicity Agent	Dehydrate Sorbitol				
Solvent	Sterile Water for injection				

Example 1F

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IV Acetaminophen Formulations

Example 1F is prepared according to the procedure outlined in Example 2 using the amounts of the excipients described in Example 1.

Excipients:	Formula 1(F)				
	Acetaminophen				
	0.550 g	0.650 g	0.750 g	0.850 g	0.950 g
Antioxidant	metabisulfite	metabisulfite	metabisulfite	metabisulfite	metabisulfite
pH Modulator	Sodium hydroxide				
pH Modulator	Hydrochloric Acid				

US 9,399,012 B2

19

-continued

20

Formula 1(F)					
Acetaminophen					
Excipients:	0.550 g	0.650 g	0.750 g	0.850 g	0.950 g
Buffering Agent	Disodium Phosphate				
Isotonicity Agent	Dehydrate Glucose				
Solvent	Sterile Water for injection				

Example 2

Preparation of IV Formulation Solutions

Prior to storage the formulations set forth in Example 1 are subjected to bubbling with nitrogen, transferred to Type II colorless bottles, and then placed under vacuum (low pressure approx. 550 mm of Hg) before stoppering the bottles with a synthetic elastomer grey stopper crimped with an aluminum cap. The residual oxygen content is approximately 1.5 ppm of dissolved oxygen. The bottles are then sterilized at 121° C. for 15 minutes. Sterile solutions are stored at ambient temperature (less than 30° C.) for up to two years prior to use.

Example 3

A Phase III Randomized, Double-Blind, Placebo-Controlled, Multi-Center, Parallel-Group, Repeated-Dose Study of the Analgesic Efficacy and Safety of 650 mg IV Acetaminophen Versus Placebo for the Treatment of Postoperative Pain After Abdominal Laparoscopic Surgery

In an effort to provide an intravenous, non-NSAID, non-opioid treatment for pain relief, the safety and efficacy of a 650 mg IV dose of APAP for the treatment of acute pain is examined.

Study Design and Evaluation

A Phase III, randomized, double-blind, Placebo-controlled, multi-center, parallel-group, repeated dose study is conducted in approximately 240 Subjects who have undergone planned or elective abdominal laparoscopic surgery. Approximately 15 to 20 US sites will participate in the Study.

Subjects will be centrally randomized, across all study centers, to receive infusions of Study Medication (either APAP or Placebo) at a dose at a dose (650 mg, 1000 mg, or placebo) and schedule described below.

Timed PI and pain relief (PR) Assessments will begin at baseline just prior to T0, the start of the first infusion of Study Medication, and continue through T24 hours.

All Subjects have access to rescue medication at all times throughout the study, as described below.

The Study will include the following assessment periods and procedures:

Screening (Day -21 to Randomization)

Screening is the period that begins when the Subject signs the Informed Consent Form and ends with randomization to Study Medication on POD1. During this period, the eligibility and baseline status of the Subject are determined.

Treatment Period (Dose 1/T0/POD1 to T24/POD2)

Administration of Study Medication (and Study-related assessments) will occur from T0 (morning of POD1) to T24 hours (morning of POD2).

15 Criteria for Evaluation

The primary efficacy endpoint is SPID24 (defined as the Sum of VAS score differences from baseline at T0 to T24), excluding all data after rescue medication.

Subject Selection Criteria

20 To be eligible for entry into the Study, Subjects must meet all of the following criteria prior to surgery: (1) Provide written Informed Consent prior to participation in the Study; (2) is scheduled to undergo abdominal laparoscopic surgery (laparoscopic gastric bypass procedures are not eligible); (3) If Subject is a female of childbearing potential, have a negative pregnancy test within 21 days of surgery; (4) be at least 18, but not more than 80 years of age; (5) Have a Body Mass Index (BMI) ≥ 19 and ≤ 45 lb/in²; (6) Have an ASA risk class of I, II, or III according to the American Society of Anesthesiologists; (7) Have the ability to read and understand the Study procedures and the use of the pain scales and have the ability to communicate meaningfully with the Study Investigator and staff; (8) Be free of other physical, mental, or medical conditions which, in the opinion of the Investigator, makes 30 Study participation inadvisable

35 Exclusion Criteria (Screening)

A Subject is NOT eligible for entry if ANY of the following criteria are met: (1) Used opioids or tramadol daily for greater than 7 days prior to Study Medication administration (Subjects who, in the Investigator's opinion have or are developing opioid tolerance are to be excluded); (2) Has been treated with Chapparal, Comfrey, Germanander, Gin Bu Huan, Kava, Pennyroyal, Skullcap, St. John's Wort, or Valerian within 14 days prior to surgery; (3) Has significant medical disease(s), laboratory abnormalities or condition(s) that in the Investigator's judgment could compromise the Subject's welfare, ability to communicate with the Study staff, complete Study activities, or would otherwise contraindicate Study participation; (4) Has known hypersensitivity to opioids, acetaminophen, or the inactive ingredients (excipients) of the Study Medication; (5) Has known or suspected history of alcohol or drug abuse or dependence within the previous 2 years; (6) Has impaired liver function, e.g., AST/ALT/bilirubin greater than or equal to 3.0 times the upper limit of normal, active hepatic disease, 45 evidence of clinically significant liver disease, or other condition (e.g., alcoholism, cirrhosis, or hepatitis) that may suggest the potential for an increased susceptibility to hepatic toxicity with Study Medication exposure; (7) Has been treated with monoamine oxidase inhibitors (MAOIs) within 7 days prior to surgery; (8) Has participated in another clinical Study (investigational or marketed product) within 30 days of surgery

55 Post Operative Exclusion Criteria

The Subject must not meet any of the following criteria 60 after surgery and prior to randomization to Study Medication: (1) Had any other surgery than the planned laparoscopic surgery or had intra operative or post operative complications

US 9,399,012 B2

21

which in the view of the Investigator would make Study participation inadvisable; (2) Has taken non steroidal anti-inflammatory drugs (NSAIDs), steroids or MAOIs during the day after surgery. Exceptions: The use of low-dose aspirin, e.g., 81 mg/day, for cardioprophylaxis, and topical or inhaled steroids are acceptable; (3) Had any neuraxial opioids or continuous local anesthetic infusions via percutaneous catheters administered as part of the anesthetic or post operative analgesic management (local anesthetic infiltration of surgical wounds at the time of closure is acceptable if done as a single injection); (4) Had a fever (greater than 38.6° C. or 101.5° F.) requiring treatment.

Postoperative Assessment (POD0)

The Subject will undergo abdominal laparoscopic surgery or other approved surgical procedure as described herein. Details of the surgical procedure(s) will be recorded on the CRF including the type of procedure(s) performed and perioperative medication will be recorded.

Example 4

Phase III, Open-Label, Prospective, Multi-Center, Repeated Dose, Randomized, Multi-Day Safety and Efficacy Study of 650 mg IV Acetaminophen

A Phase III, open-label, prospective, multi-center, repeated dose, randomized, multi-day safety and efficacy study was conducted in 213 subjects. The subjects were randomized as follows: 92 subjects to a q6 group (1 g of IV acetaminophen every 6 hours), 91 subjects to a q4 group (650 mg of IV acetaminophen every 4 hours), and 28 subjects to a standard of care control group, which could include oral acetaminophen, but no IV acetaminophen. Subjects who completed 5 days of study treatment included 63 in the q6 group, 59 in the q4 group and 26 in the control group. The primary endpoint was an assessment of safety using spontaneous adverse event reporting and daily liver enzymes. Efficacy evaluations were also performed.

Inclusion Criteria (Screening)

To be eligible for entry into the Study, Subjects had to meet all the following criteria: (1) Provide written informed consent prior to participation in the Study; (2) Be at least 18 years of age and weigh at least 41 kg; (3) Be anticipated by the Investigator to require multi-day (target is five days) use of IV treatment either because of: (a) having a “nothing by mouth” (NPO) status, (b) having a medical condition that makes oral intake difficult, or (c) having a medical condition that requires IV treatment; (4) Be willing to undergo 5 days of treatment with IV acetaminophen for the treatment of pain or fever (defined as a core temperature $\geq 38^{\circ}$ C.). Subjects had a slightly less than 15% chance (one in seven) of being assigned to the Control Group and receiving standard of care treatment, but no IV APAP; (5) Have the ability to read and understand the Study procedures and have the ability to communicate meaningfully with the Study Investigator and staff, and (6) If a female of child bearing potential, have a negative pregnancy test within 48 hours of randomization.

Exclusion Criteria (Screening)

A Subject was not eligible for entry if any of the following criteria were met: (1) Had a significant medical disease, laboratory abnormality or condition that, in the Investigator's judgment, could compromise the Subject's welfare or would otherwise contraindicate Study participation; (2) Was expected to have difficulty in communicating with the Study staff or completing Study requirements (including follow up visits); (3) Had known hypersensitivity to acetaminophen or the inactive ingredients (excipients) of IV acetaminophen or

22

any contraindication to receiving acetaminophen; (4) Had impaired liver function, e.g., ALT greater than or equal to 3 times the upper limit of normal (ULN), bilirubin greater than or equal to 3 times ULN, known active hepatic disease (e.g., hepatitis), evidence of clinically significant chronic liver disease or other condition affecting the liver (e.g., alcoholism as defined by DSM-IV, cirrhosis or chronic hepatitis); or (5) Had participated in an interventional clinical Study (investigational or marketed product) within 30 days of Study entry.

10 Efficacy Analysis

All analyses of efficacy were conducted on the modified intent-to-treat population separately for the two indications (acute pain and fever). Subjects' Global Evaluations were summarized descriptively (m, mean, SD, median, minimum, and maximum) by treatment group for each study day and for overall assessments. Summary statistics were also provided for each site.

15 Comparisons of efficacy endpoints between the following

pairs of treatment groups were investigated using two-sided tests at the 5% level of significance:

IV acetaminophen 1 g versus IV acetaminophen 650 mg
IV acetaminophen 1 g versus standard of care treatment
IV acetaminophen 650 mg versus standard of care treatment

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A one-way analysis of variance (ANOVA) model with treatment group as the factor was used to test the treatment difference between these pairs. All groups were included in this analysis model. The p-values from the ANOVA model were presented along with the summary statistics.

Safety Analyses

All analyses of safety were conducted on the safety population.

Percentage of subjects withdrawn due to adverse event, percentage of subjects with adverse events (AEs) or serious adverse events (SAEs), and percentage of subjects with clinically meaningful changes in laboratory parameters were summarized.

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All adverse events and serious adverse events were coded according to the Medical Dictionary for Regulatory Activities (MedDRA), Version 10.0. Additional analyses included displays of the number of subjects reporting at least one AE (incidence table), total number of episodes of each AE by body system and by severity, total number of episodes of each AE by body system, and by attribution. Liver function test abnormalities were graded using the Common Terminology Criteria for Adverse Events.

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For each clinical laboratory parameter, descriptive statistics (m, mean, standard deviation, median, and range) were tabulated for baseline and final values. Change from baseline was tabulated for those subjects who had both baseline and final values. Liver function tests were also evaluated using values that were normalized to the upper limit of normal values for the local laboratory.

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A shift table was prepared to present the shift in baseline clinical laboratory values that were clinically relevantly high or low at baseline and/or final measurement.

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Descriptive statistics (m, mean, standard deviation, median, and range) were tabulated for changes in vital signs from baseline to final measurement.

Results

Disposition of Subjects

A total of 257 subjects were screened for study enrollment. 60 Of the total screened, 44 were screen failures, and 213 were enrolled and randomized: 92 subjects in the q6 group, 91 subjects in the q4 group, and 28 subjects in the control group.

US 9,399,012 B2

23

Subjects who completed 5 days of study treatment included 63 in the q6 group, 59 in the q4 group and 26 in the control group.

Subjects in the q4h group and q6h group were considered to be a Study Treatment Discontinuation/Early Termination if they received at least one dose of IV acetaminophen and discontinued study participation prior to completion of Day 5 treatments. Subjects in the control group were considered to be a Study Treatment Discontinuation/Early Termination if they discontinued at any time after T0, but prior to completion of Day 5 standard of care treatments.

Subjects who received at least one dose of IV acetaminophen and discontinued study participation prior to completing Day 5 treatments, but returned for the Last Study Visit were considered as "Partial Treatment Completers". Similarly, subjects in the control group who discontinued study participation prior to completing Day 5 standard of care treatments, but returned for the Last Study Visit were considered as "Partial Treatment Completers".

Subjects who completed Day 5 treatments (IV acetaminophen or standard of care) and procedures were characterized as a "Treatment Completer". A Treatment Completer who elected to discontinue study participation prior to the Last Study Visit was characterized as a Treatment Completer Early Termination.

Safety Outcome

There were no clinically relevant differences between the treatment groups in the frequency of serious, severe, related, or overall treatment emergent adverse events (TEAEs). In fact, most TEAEs were assessed by the Investigator to be mild or moderate in severity. The frequency of liver enzyme elevations seen in the treatment groups was comparable. More specifically, with regard to the hepatic transaminases alanine aminotransferase and aspartate aminotransferase, the frequency and severity of the elevations were comparable between the treatment groups. There were no clinically relevant differences between the treatment groups regarding laboratory assessments, vital signs, or physical examinations. Thus, based on these data, intravenous acetaminophen in both active treatment groups (i.e., 650 mg and 1000 mg dose groups) was well tolerated.

Efficacy Outcome

The modified intent-to-treat population was used for all analyses of efficacy: Subject Global Evaluations (rating of study treatments and rating of satisfaction with side effects related to study treatments) provided as a daily lookback (days 2 through 5) and overall evaluation (overall treatment period lookback) using a 4 point categorical rating scale (0=poor, 1=fair, 2=good, 3=excellent). A one-way ANOVA model with treatment group as the factor was used to test the treatment difference between each treatment pair:

IV acetaminophen 1 g q6h versus IV acetaminophen 650 mg q4h

IV acetaminophen 1 g q6h versus standard of care treatment (Control)

IV acetaminophen 650 mg q4h versus standard of care treatment (Control)

All endpoints were tested at the 0.05 significance level (two-sided).

The IV acetaminophen 650 mg q4h group relative to the control group produced statistically significantly better satisfaction ratings for the Subject Global Assessments rating the level of satisfaction with the side effects related to study treatments the on the day 5 (mean rating 2.4 vs. 2.0, p=0.0167) and at the end of day 5 prior to discharge (mean rating 2.4 vs. 2.0, p=0.0129) 24 h look back assessments. On day 4, the satisfaction rating showed a trend to significance

24

(mean rating 2.5 vs. 2.2, p=0.1162). With respect to the Subject Global Assessments rating the level of satisfaction with the study treatments, there were no statistically significant differences between the IV acetaminophen 650 mg q6h group and control group at any of the assessment points. For the both of the Subject Global Assessments rating either the level of satisfaction with the study treatments or the level of satisfaction with the side effects related to study treatments, there was no statistically significant differences between the two active treatment groups with respect to the daily 24 h lookback assessments on day 2, day 3, day 4, day 5, or at the end of day 5 prior to discharge; nor was there a statistically significant difference on the overall assessment at the Study Completion Visit.

The IV acetaminophen 1 g q6h group produced statistically significantly better satisfaction ratings for the Subject Global Assessments rating the level of satisfaction with the side effects related to study treatments on day 5 (mean rating 2.4 vs. 2.0, p=0.0062) and at the end of day 5 prior to discharge (mean rating 2.5 vs. 2.0, p=0.0073) 24 h lookback assessments compared to the control group. On day 4, the satisfaction rating showed a trend to significance (mean rating 2.5 vs. 2.2, p=0.0744). With respect to the Subject Global Assessments rating the level of satisfaction with the study treatments, there were no statistically significant differences between the IV acetaminophen 1 g q6h group and control group at any of the assessment points.

Statistically significant differences were observed for both active treatment groups versus the control group in the Subject Global Assessments rating the level of Satisfaction with the side effects related to study treatments on the day 5 and on the end of day 5 prior to discharge daily 24 h lookback assessments. Thus, these data suggest that the IV acetaminophen 1 g q6h and 650 mg q4h groups were efficacious and provided comparable efficacy based upon the global satisfaction ratings.

Many modifications, equivalents, and variations of the present invention are possible in light of the above teachings, therefore, it is to be understood that within the scope of the appended claims, the invention may be practiced other than as specifically described.

What is claimed is:

1. A method for the treatment of pain or fever in an adult human or an adolescent human subject weighing at least 50 kg, in need thereof, comprising administering to the subject, by an intravenous route of administration, a therapeutically effective amount of a pharmaceutical composition comprising about 550 mg to about 800 mg of acetaminophen; and repeating said administration at least once at an interval of about 3 to about 5 hours.

2. The method of claim 1, wherein the subject receives a total of about 3 to about 5 grams of acetaminophen in a period of twenty four hours.

3. The method of claim 2, wherein the pharmaceutical composition is administered at least six times in a period of twenty four hours.

4. The method of claim 1, wherein the pharmaceutical composition comprises at least one antioxidant.

5. The method of claim 4, wherein the at least one antioxidant is selected from the group consisting of cysteine hydrochloride monohydrate, thioglycolic acid, thiolacetic acid, dithiothreitol, reduced glutathione, thiourea, a-thioglycerol, cysteine, acetylcysteine, mercaptoethane sulfonic acid, ascorbic acid, ascorbic acid derivatives, an organic compound having at least one thiol, an alkyl polyhydroxylated compound, a cycloalkyl polyhydroxylated compound, and mixtures thereof.

US 9,399,012 B2

25

6. The method of claim 5, wherein the at least one antioxidant comprises cysteine hydrochloride monohydrate.

7. The method of claim 1, further comprising a buffering agent.

8. The method of claim 7, wherein the buffering agent comprises disodium phosphate dehydrate.

9. The method of claim 1, wherein the pharmaceutical composition has a pH between about 4 to about 8.

10. The method of claim 9, wherein the pharmaceutical composition has a pH of between about 5 and about 6.

11. The method of claim 1, wherein the pharmaceutical composition has an osmolality of between about 200 mOsm/L to about 400 mOsm/L.

12. The pharmaceutical composition of claim 11, wherein the acetaminophen is present in the composition in an amount of about 600 mg to about 700 mg.

13. The method of claim 1, wherein the pharmaceutical composition further comprises an isotonicity agent.

14. The method of claim 13, wherein the isotonicity agent is dextrose, mannitol, or lactose.

15. The method of claim 14, wherein the isotonicity agent is mannitol.

16. The method of claim 1, further comprising EDTA.

17. The method of claim 1, wherein the pharmaceutical composition further comprises at least one analgesic agent other than acetaminophen.

18. The method of claim 17, wherein the at least one analgesic agent other than acetaminophen comprises an anilide, an opioid, an NSAID, a cannabinoid, a pyrazalone, or a barbiturate.

19. The method of claim 18, wherein the at least one analgesic agent other than acetaminophen comprises an opioid.

20. The method of claim 1, wherein the subject is suffering from a fever.

21. The method of claim 1, wherein the subject is unconscious, sedated, fasting, nauseous, or unable to be administered a pharmaceutical composition by an oral route.

22. The method of claim 1, wherein the pharmaceutical composition is administered after a surgical intervention on the subject.

23. The method of claim 1, wherein the pharmaceutical composition is administered within 3 hours of a surgical intervention on the subject.

24. The method of claim 23, wherein the pharmaceutical composition is administered within 1 hour of a surgical intervention on the subject.

25. The method of claim 1, wherein the pharmaceutical composition is administered postoperatively.

26. The method of claim 1, further comprising administering to the subject at least one analgesic agent other than acetaminophen.

27. The method of claim 26, wherein the at least one analgesic agent other than acetaminophen comprises an anilide, an opioid, an NSAID, a cannabinoid, a pyrazalone, or a barbiturate.

28. The method of claim 1, wherein the subject is suffering from an infection.

29. A method for reducing pain or fever in an adult human or an adolescent human subject weighing at least 50 kg, in need thereof, comprising administering to the subject, by an intravenous route of administration, a therapeutically effective amount of a pharmaceutical composition in solution comprising: about 600 mg to about 700 mg of acetaminophen, cysteine hydrochloride monohydrate, disodium phosphate dehydrate, and mannitol, wherein the solution has a pH of about 5 to about 6, and an osmolality of about 200-400

26

mOsm/L; and repeating said administration at least once at an interval of about 3 to about 5 hours.

30. The method of claim 29, wherein the pharmaceutical composition has an acetaminophen concentration of about 0.5% (w/v) to about 10% (w/v).

31. The method of claim 30, wherein the acetaminophen concentration is about 1% (w/v).

32. The method of claim 29, wherein the pharmaceutical composition further comprises EDTA.

33. The method of claim 29, wherein the subject is suffering from postoperative pain.

34. The method of claim 1, wherein the level of pain the subject is suffering from is reduced.

35. The method of claim 1, wherein the pharmaceutical composition is administered as a pretreatment.

36. The method of claim 29, wherein the administered dose of acetaminophen is 650 mg, and further comprising repeating intravenous administration of 650 mg acetaminophen at least once at an interval of about 3 hours to about 5 hours.

37. The method of claim 36, wherein the interval is about 4 hours.

38. The method of claim 1, further comprising repeating intravenous administration of about 600 mg to about 700 mg acetaminophen at least once at an interval of about 3 hours to about 5 hours.

39. The method of claim 1, wherein the administered dose of acetaminophen is 650 mg, and further comprising repeating intravenous administration of 650 mg acetaminophen at least once at an interval of about 3 hours to about 5 hours.

40. The method of claim 29, wherein the interval is about 4 hours.

41. The method of claim 1, wherein the composition may be administered to the subject without dilution.

42. The method of claim 1, wherein the composition is a sterile solution that is ready for direct administration to the subject.

43. The method of claim 1, wherein the pharmaceutical composition is a lyophilized powder.

44. The method of claim 43, wherein the lyophilized powder must be reconstituted in solution prior to administration.

45. The method of claim 1, wherein the pharmaceutical composition is intravenously administered to the subject over about 5 minutes to about 30 minutes.

46. The method of claim 45, wherein the pharmaceutical composition is intravenously administered to the subject over about 15 minutes.

47. The method of claim 1, wherein the subject is administered less than 4 grams of acetaminophen over a twenty-four hour period.

48. A method of treating pain in a human subject weighing at least 50 kg comprising:

administering to the subject a therapeutically effective amount of a pharmaceutical composition comprising about 600 mg to about 700 mg of acetaminophen; and repeating said administration at least about once every 4 hours;

wherein the composition is administered to the subject as a 15-minute intravenous infusion, and wherein the subject is administered less than 4 grams of acetaminophen over a 24-hour period.

49. The method of claim 48, wherein the composition comprises about 650 mg of acetaminophen.

50. The method of claim 48, wherein the composition is administered to the subject at least six times in a period of 24 hours.

US 9,399,012 B2

27

51. The method of claim 48, wherein the composition is administered to the subject at least seven times in a period of 24 hours.

52. The method of claim 48, wherein the composition is administered to the subject at least eight times in a period of 24 hours.

53. The method of claim 48, wherein the composition may be administered to the subject without dilution.

54. The method of claim 48, wherein the composition is a sterile solution that is ready for subject administration.

55. The method of claim 48, wherein the composition is a lyophilized powder.

56. The method of claim 55, wherein the lyophilized powder must be reconstituted in solution prior to subject administration.

57. The method of claim 48, wherein the composition may be stored at ambient temperature for two years prior to use.

58. The method of claim 57, wherein the ambient temperature is less than 30 degrees Celsius.

59. The method of claim 48, wherein the composition further comprises a buffer in an amount of about 0.1% to about 0.7%.

60. The method of claim 48, wherein the composition further comprises a buffer in an amount of about 0.2% to about 0.3%.

61. The method of claim 48, wherein the composition further comprises a buffer in an amount of about 0.05% to about 2.0%.

62. The method of claim 48, wherein the composition further comprises at least one buffering agent.

63. The method of claim 62, wherein the at least one buffering agent is selected from the group consisting of a pharmaceutically acceptable salt or acid of citrate, phosphate, acetate, glutamate, tartrate, benzoate, lactate, histidine or other amino acids, gluconate, malate, succinate, formate, propionate, carbonate, or any combination thereof.

64. The method of claim 63, wherein the at least one buffering agent is disodium phosphate dehydrate.

65. The method of claim 48, wherein the pharmaceutical composition has a pH between about 4 to about 8.

66. The method of claim 48, wherein the pharmaceutical composition has a pH of between about 5 and about 6.

67. The method of claim 48, wherein the composition further comprises an antioxidant in an amount of about 0.3% to about 1.0%.

68. The method of claim 48, wherein the composition further comprises an antioxidant in an amount of about 0.5%.

69. The method of claim 48, wherein the composition further comprises at least one antioxidant.

70. The method of claim 69, wherein the at least one antioxidant is selected from the group consisting of cysteine hydrochloride monohydrate, thioglycolic acid, thiolacetic acid, dithiothreitol, reduced glutathione, thiourea, a-thioglycerol, cysteine, acetylcysteine, mannitol, mercaptoethane sulfonic acid, ascorbic acid, ascorbic acid derivatives, an organic compound having at least one thiol, an alkyl polyhydroxylated compound, a cycloalkyl polyhydroxylated compound, and mixtures thereof.

71. The method of claim 70, wherein the at least one antioxidant is cysteine hydrochloride monohydrate.

72. The method of claim 70, wherein the at least one antioxidant is mannitol.

73. The method of claim 48, wherein the composition further comprises an isotonicity agent in an amount of about 65% to about 85%.

74. The method of claim 48, wherein the composition further comprises at least one isotonicity agent.

28

75. The method of claim 74, wherein the at least one isotonicity agent is selected from the group consisting of mannitol, sorbitol, glycerol, sucrose, glucose, dextrose, levulose, fructose, lactose, polyethylene glycols 400 to 4000, phosphates, sodium chloride, potassium chloride, calcium chloride, calcium gluconoglucoheptonate, dimethyl sulfone, and combinations thereof.

76. The method of claim 75, wherein the at least one isotonicity agent is mannitol.

77. The method of claim 48, wherein the composition has an osmolality of between about 200 mOsm/L to about 400 mOsm/L.

78. The method of claim 48, wherein the composition comprises 650 mg of acetaminophen, and wherein the subject is administered the composition every four hours in a twenty-four hour period.

79. A method of reducing fever in a human subject weighing at least 50 kg comprising:

administering to the subject a therapeutically effective amount of a pharmaceutical composition comprising about 600 mg to about 700 mg of acetaminophen; and repeating said administration at least about once every 4 hours;

wherein the composition is administered to the subject as a 15-minute intravenous infusion, and

wherein the subject is administered less than 4 grams of acetaminophen over a twenty-four hour period.

80. The method of claim 79, wherein the composition comprises about 650 mg of acetaminophen.

81. The method of claim 79, wherein the composition is administered to the subject at least six times in a period of 24 hours.

82. The method of claim 79, wherein the composition is administered to the subject at least seven times in a period of 24 hours.

83. The method of claim 79, wherein the composition is administered to the subject at least eight times in a period of 24 hours.

84. The method of claim 79, wherein the composition may be administered to the subject without dilution.

85. The method of claim 79, wherein the composition is a sterile solution that is ready for subject administration.

86. The method of claim 79, wherein the composition is a lyophilized powder.

87. The method of claim 86, wherein the lyophilized powder must be reconstituted in solution prior to administration.

88. The method of claim 79, wherein the composition may be stored at ambient temperature for two years prior to use.

89. The method of claim 88, wherein the ambient temperature is less than 30 degrees Celsius.

90. The method of claim 79, wherein the composition further comprises a buffer in an amount of about 0.1% to about 0.7%.

91. The method of claim 79, wherein the composition further comprises a buffer in an amount of about 0.2% to about 0.3%.

92. The method of claim 79, wherein the composition further comprises a buffer in an amount of about 0.05% to about 2.0%.

93. The method of claim 79, wherein the composition further comprises at least one buffering agent.

94. The method of claim 93, wherein the at least one buffering agent is selected from the group consisting of a pharmaceutically acceptable salt or acid of citrate, phosphate, acetate, glutamate, tartrate, benzoate, lactate, histidine or other amino acids, gluconate, malate, succinate, formate, propionate, carbonate, or any combination thereof.

60 65

US 9,399,012 B2

29

95. The method of claim **94**, wherein the at least one buffering agent is disodium phosphate dehydrate.

96. The method of claim **79**, wherein the pharmaceutical composition has a pH between about 4 to about 8.

97. The method of claim **79**, wherein the pharmaceutical composition has a pH of between about 5 and about 6.

98. The method of claim **79**, wherein the composition further comprises an antioxidant in an amount of about 0.3% to about 1.0%.

99. The method of claim **79**, wherein the composition further comprises an antioxidant in an amount of about 0.5%.

100. The method of claim **79**, wherein the composition further comprises at least one antioxidant.

101. The method of claim **100**, wherein the at least one antioxidant is selected from the group consisting of cysteine hydrochloride monohydrate, thioglycolic acid, thiolacetic acid, dithiothreitol, reduced glutathione, thiourea, α -thioglycerol, cysteine, acetylcysteine, mannitol, mercaptoethane sulfonic acid, ascorbic acid, ascorbic acid derivatives, an organic compound having at least one thiol, an alkyl polyhydroxylated compound, a cycloalkyl polyhydroxylated compound, and mixtures thereof.

102. The method of claim **101**, wherein the at least one antioxidant is cysteine hydrochloride monohydrate.

30

103. The method of claim **101**, wherein the at least one antioxidant is mannitol.

104. The method of claim **79**, wherein the composition further comprises an isotonicity agent in an amount of about 65% to about 85%.

105. The method of claim **79**, wherein the composition further comprises at least one isotonicity agent.

106. The method of claim **105**, wherein the at least one isotonicity agent is selected from the group consisting of ¹⁰mannitol, sorbitol, glycerol, sucrose, glucose, dextrose, levulose, fructose, lactose, polyethylene glycols 400 to 4000, phosphates, sodium chloride, potassium chloride, calcium chloride, calcium gluconoglucoheptonate, dimethyl sulfone, and combinations thereof.

107. The method of claim **106**, wherein the at least one isotonicity agent is mannitol.

108. The method of claim **79**, wherein the composition has an osmolality of between about 200 mOsm/L to about 400 mOsm/L.

109. The method of claim **79**, wherein the pharmaceutical composition comprises 650 mg of acetaminophen, and wherein the subject is administered the composition every four hours in a twenty-four hour period.

* * * * *

EXHIBIT B

(12) **United States Patent**
Royal et al.

(10) **Patent No.:** **US 9,610,265 B2**
(45) **Date of Patent:** ***Apr. 4, 2017**

(54) **REDUCED DOSE INTRAVENOUS ACETAMINOPHEN**

(71) Applicant: **MALLINCKRODT IP**, Dublin (IE)

(72) Inventors: **Mike Allan Royal**, San Diego, CA (US); **James Bradley Breitmeyer**, Rancho Santa Fe, CA (US)

(73) Assignee: **MALLINCKRODT IP**, Damastown Industrial Estate, Dublin (IE)

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(58) **Field of Classification Search**

USPC 514/629
See application file for complete search history.

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Primary Examiner — San-Ming Hui

(74) *Attorney, Agent, or Firm* — Mayer Brown LLP

(57) **ABSTRACT**

Described herein are compositions and methods for intravenous administration of acetaminophen at a single dose level of less than about 1000 mg for the treatment or prevention of pain (e.g., postoperative pain) and/or fever.

20 Claims, No Drawings

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Page 2

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US 9,610,265 B2

1

REDUCED DOSE INTRAVENOUS ACETAMINOPHEN**CROSS-REFERENCE TO RELATED APPLICATIONS**

This application is a continuation of U.S. patent application Ser. No. 12/270,796, filed on Nov. 13, 2008, which is related to and claims priority to U.S. Provisional Patent Application No. 60/987,761, entitled "Reduced Dose Intravenous Acetaminophen" filed on Nov. 13, 2007, which are incorporated herein by reference in their entirety to the full extent permitted by law.

BACKGROUND OF THE INVENTION

In the hospital, particularly in the postoperative setting, pain is a primary concern of patients. Opioid analgesics have been used to treat postoperative pain since 1784 and parenteral morphine has been a primary treatment modality since the 1850s. While opioids are highly effective in the treatment of many painful conditions, they have side effects and dose-dependent risks including nausea, vomiting, constipation, urinary retention, sedation, and respiratory depression. Similarly, non steroidal anti-inflammatory drugs (NSAIDs), including the older non selective (dual inhibitor) products and newer cyclo-oxygenase (COX)-2 products, have a variety of unwanted side effects especially when used in the perioperative setting. Non selective NSAIDs are associated with platelet dysfunction and the potential for bleeding at the surgical site, upper gastrointestinal ulcers and bleeding, edema, hypertension, congestive heart failure, renal dysfunction, severe skin reactions such as Stevens-Johnson syndrome and toxic epidermal necrolysis, anaphylaxis, and most recently, an increased risk of thrombotic cardiovascular events.

SUMMARY OF THE INVENTION

Described herein are pharmaceutical compositions having a reduced dose of acetaminophen for intravenous administration, and methods of using these compositions for treating and/or preventing pain and/or fever in a subject.

In some embodiments, the pharmaceutical compositions described herein comprise less than about 1 gram of acetaminophen, wherein the pharmaceutical composition is provided as a formulation suitable for intravenous administration. For example, various embodiments may comprise about 500 mgs to about 1 gram, or about 500 mgs to about 800 mgs, or about 500 mgs to about 750 mgs. In various embodiments, the pharmaceutical compositions described herein comprise about 600 mg to about 700 mg of acetaminophen.

In some embodiments, the pharmaceutical compositions described herein further comprise at least one antioxidant. In some embodiments, the at least one antioxidant comprises cysteine hydrochloride monohydrate, thioglycolic acid, thiolacetic acid, dithiothreitol, reduced glutathione, thiourea, alpha-thioglycerol, cysteine, acetylcysteine, or mercaptoethane sulfonic acid, ascorbic acid ascorbic acid derivatives, an organic compound having at least one thiol, an alkyl polyhydroxylated compound, or a cycloalkyl polyhydroxylated compound.

In some embodiments, the pharmaceutical composition further comprises a buffering agent (e.g., disodium phosphate dehydrate). In some embodiments, the pharmaceutical composition has a pH from about 4 to about 8 when in

2

solution. In some embodiments, the pharmaceutical composition has a pH of about 5 to about 6 when in solution.

In some embodiments, the pharmaceutical composition has an osmolality from about 250 mOsm/L to about 400 mOsm/L when in solution. In some embodiments, the pharmaceutical composition further comprises an isotonicity agent. In some embodiments, the isotonicity agent is dextrose, mannitol, or lactose.

In some embodiments, the pharmaceutical composition further comprises at least one analgesic agent other than acetaminophen. In some embodiments, the at least one analgesic agent other than acetaminophen comprises an anilide, an opioid, an NSAID, a cannabinoid, a pyralazone, or a barbiturate.

In some embodiments, the pharmaceutical composition further comprises EDTA.

In a further aspect provided herein is a method for preventing or reducing pain or fever in a subject in need thereof, comprising administering to the subject, by an intravenous route of administration, a pharmaceutical composition described herein. In some embodiments, the administration is repeated at least once with an interval of about 3 to about 5 hours. In some embodiments, the administration is repeated at least six times in a period of twenty four hours. In various embodiments, the administration is repeated three to eight times (e.g., 3 times, 4 times, five times, six times, seven times, or eight times) in a period of twenty four hours and about 3 to about 5 grams of acetaminophen (e.g., about 3 grams, about 4 grams or about 5 grams) is delivered over the twenty four hour period. In other embodiments, the administration is repeated three to eight times in a period of twenty four hours and less than about 4 grams of acetaminophen is delivered over the twenty four hour period.

In some embodiments, the pharmaceutical formulation for IV administration is a solution comprising about 600 mg to about 700 mg of acetaminophen, cysteine hydrochloride monohydrate, disodium phosphate dehydrate, and mannitol, wherein the solution has a pH of between about 5 and about 6 and an osmolality of between about 200-400 mOsm/L. In some embodiments, the pharmaceutical composition in solution has an acetaminophen concentration of about 0.5% (w/v) to about 10% (w/v). In some embodiments, the acetaminophen concentration is about 1% (w/v). In some embodiments, the pharmaceutical composition to be administered further comprises EDTA.

In some embodiments, the subject to be treated is suffering from an infection. In some embodiments, the subject to be treated is suffering from a fever. In some embodiments, the subject to be treated is unconscious, sedated, fasting, nauseous, or unable to be administered a pharmaceutical composition by an oral route.

In some embodiments, the pharmaceutical composition is administered to the subject after a surgical intervention. In some embodiments, the pharmaceutical composition is administered within three hours of a surgical intervention on the subject. In some embodiments, the pharmaceutical composition is administered within 1 hour of a surgical intervention on the subject. In some embodiments, the pharmaceutical composition is administered postoperatively. In some embodiments, the subject to be treated is suffering from postoperative pain.

In various embodiments the pharmaceutical compositions described herein are administered as a pretreatment.

In another aspect provided herein is a method for preventing or reducing pain or fever in a subject in need thereof,

US 9,610,265 B2

3

comprising administering to the subject, by an intravenous route of administration, a pharmaceutical composition described herein.

INCORPORATION BY REFERENCE

All publications and patent applications mentioned in this specification are herein incorporated by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference.

DETAILED DESCRIPTION OF THE INVENTION

To date, the standard intravenous (IV) dose of acetaminophen for the relief of pain or fever has been 1000 mg in adults and adolescents weighing at least 50 kg. At this dose level, the frequency of acetaminophen administration is limited to a maximum of once every six hours (i.e., four administrations per twenty four hours) to minimize the potential for hepatotoxicity. On the other hand, it has generally been noted that acetaminophen has greatest efficacy during its initial rise in plasma concentration, i.e., during the first few hours post-administration, and is less effective later on after the plasma concentration of the drug drops from its peak. While not wishing to be bound by theory, it is thought that this change in efficacy is likely due to a time and concentration-dependent modulation of the central and peripheral nociceptive pathways through which acetaminophen acts.

Further, if the duration of effect of a 1000 mg dose of acetaminophen is shorter in duration than 6 hours, the use of this dose is limited since dosing more frequently than every 6 hours, e.g., every 4 hours, leaves a gap in coverage due to the 4 g acetaminophen maximum daily limit. In the treatment of fever, a dose less than 1000 mg may be fully effective due to the fact that a lower plasma level (compared to that needed for pain) is needed to effectively reduce fever.

Thus, intravenous administration of a reduced dose of acetaminophen, as described herein, permits more frequent IV acetaminophen administration to yield better overall relief of symptoms for many patients while avoiding any potential gap artificially created by the daily limit.

Also, the reduced acetaminophen IV dose affords greater flexibility to the physician in customizing treatments to the needs of the patient, selecting the dose of other drugs for use in combination therapies and allowing for smoother transitions to oral products containing acetaminophen.

Accordingly, described herein are reduced IV dose formulations of acetaminophen for intravenous administration and the use of reduced IV doses of acetaminophen for use for the treatment or prevention of pain and/or fever.

Certain Terminology

Unless defined otherwise, all technical and scientific terms used herein have the same meaning as is commonly understood by one of skill in the art to which the claimed subject matter belongs. In the event that there is a plurality of definitions for terms herein, those in this section prevail. Where reference is made to a URL or other such identifier or address, it is understood that such identifiers can change and particular information on the internet can come and go, but equivalent information can be found by searching the internet or other appropriate reference source. Reference thereto evidences the availability and public dissemination of such information.

4

It is to be understood that the foregoing general description and the following detailed description are exemplary and explanatory only and are not restrictive of any subject matter claimed. In this application, the use of the singular includes the plural unless specifically stated otherwise. It must be noted that, as used in the specification and the appended claims, the singular forms "a", "an" and "the" include plural referents unless the context clearly dictates otherwise. It should also be noted that use of "or" means "and/or" unless stated otherwise. Furthermore, use of the term "including" as well as other forms, such as "include", "includes", and "included" is not limiting.

Definition of standard chemistry terms may be found in reference works, including Carey and Sundberg "ADVANCED 15 ORGANIC CHEMISTRY 4TH ED." Vols. A (2000) and B (2001), Plenum Press, New York. Unless otherwise indicated, conventional methods of mass spectroscopy, NMR, HPLC, IR and UV/Vis spectroscopy and pharmacology, within the skill of the art are employed. Unless specific definitions are 20 provided, the nomenclature employed in connection with, and the laboratory procedures and techniques of, analytical chemistry, synthetic organic chemistry, and medicinal and pharmaceutical chemistry described herein are those known in the art. Standard techniques can be used for chemical 25 syntheses, chemical analyses, pharmaceutical preparation, formulation, and delivery, and treatment of patients. Reactions and purification techniques can be performed e.g., using kits of manufacturer's specifications or as commonly accomplished in the art or as described herein. The foregoing techniques and procedures can be generally performed 30 of conventional methods well known in the art and as described in various general and more specific references that are cited and discussed throughout the present specification. Throughout the specification, groups and substituents 35 thereof can be chosen by one skilled in the field to provide stable moieties and compounds.

The terms "treat," "treating" or "treatment," and other grammatical equivalents as used herein, include alleviating, 40 abating or ameliorating a disease or condition symptoms, preventing additional symptoms, ameliorating or preventing the underlying metabolic causes of symptoms, inhibiting the disease or condition, e.g., arresting the development of the disease or condition, relieving the disease or condition, causing regression of the disease or condition, relieving a 45 condition caused by the disease or condition, or stopping the symptoms of the disease or condition, and are intended to include prophylaxis. The terms further include achieving a therapeutic benefit and/or a prophylactic benefit. By therapeutic benefit is meant eradication or amelioration of the 50 underlying disorder being treated. Also, a therapeutic benefit is achieved with the eradication or amelioration of one or more of the physiological symptoms associated with the underlying disorder such that an improvement is observed in the patient, notwithstanding that the patient may still be 55 afflicted with the underlying disorder. For prophylactic benefit, the compositions may be administered to a patient at risk of developing a particular disease, or to a patient reporting one or more.

The terms "effective amount," "therapeutically effective amount" or "pharmaceutically effective amount" as used 60 herein, refer to a sufficient amount of at least one agent or compound being administered which will relieve to some extent one or more of the symptoms of the disease or condition being treated. The result can be reduction and/or alleviation of the signs, symptoms, or causes of a disease, or 65 any other desired alteration of a biological system. For example, an "effective amount" for therapeutic uses is the

US 9,610,265 B2

5

amount of the composition comprising the compound as disclosed herein required to provide a clinically significant decrease in pain. An additional example is that an “effective amount” may be a dosage that decreases a fever. An appropriate “effective” amount in any individual case may be determined using techniques, such as a dose escalation study.

The terms “administer,” “administering,” “administration,” and the like, as used herein, refer to the methods that may be used to enable delivery of compounds or compositions to the desired site of biological action. These methods include, but are not limited to oral routes, intraduodenal routes, parenteral injection (including intravenous, subcutaneous, intraperitoneal, intramuscular, intravascular or infusion), topical and rectal administration. Those of skill in the art are familiar with administration techniques that can be employed with the compounds and methods described herein, e.g., as discussed in Goodman and Gilman, *The Pharmacological Basis of Therapeutics*, current ed.; Pergamon; and Remington’s, *Pharmaceutical Sciences* (current edition), Mack Publishing Co., Easton, Pa. In preferred embodiments, the compositions comprising acetaminophen as described herein are administered intravenously.

The term “acceptable” as used herein, with respect to a formulation, composition or ingredient, means having no persistent detrimental effect on the general health of the subject being treated.

The term “antioxidant” refers to a compound that prevents oxygen or oxygen-derived free radicals from interacting with other substances. Antioxidants are added to minimize or retard oxidative processes that occur with some drugs or excipients upon exposure to oxygen or in the presence of free radicals. These processes can often be catalyzed by light, temperature, hydrogen on concentration, presence of trace metals or peroxides.

The terms “enhance” or “enhancing,” as used herein, means to increase or prolong either in potency or duration a desired effect. Thus, in regard to enhancing the effect of therapeutic agents, the term “enhancing” refers to the ability to increase or prolong, either in potency or duration, the effect of other therapeutic agents on a system. An “enhancing-effective amount,” as used herein, refers to an amount adequate to enhance the effect of another therapeutic agent in a desired system.

“Concurrent administration,” “administered in combination” or similar phrases referring to the acetaminophen and at least one additional component means that the components are administered concurrently to the mammal being treated. By “concurrently,” it is meant that each component may be administered at the same time or sequentially in any order at different points in time. However, if not administered at the same time, they should be administered sufficiently closely in time so as to provide the desired enhancement of treatment effect. Suitable dosing intervals and the order of administration with such compounds will be readily apparent to those skilled in the art, once armed with the present disclosure. Preferably both components are administered at the same time or within the same hour.

As used herein, the term “animal” shall refer to a vertebrate animal. More preferably, the vertebrate animal is a mammal. As used herein, the term “mammal” shall refer to the Mammalia class of higher vertebrates. The term “mammal” includes, but is not limited to, a human.

As used herein, the term “pain” shall refer to all types of pain, including, but not limited, to nociceptive pain, neuropathic pain, post-operative pain, lower back pain, cluster headaches, herpes neuralgia, phantom limb pain, central

6

pain, dental pain, opioid-resistant pain, visceral pain, surgical pain, bone injury pain, pain during labor and delivery, pain resulting from burns, including sunburn, post partum pain, migraine, and genitourinary tract-related pain including cystitis. Levels of pain in a subject can be quantified using standard subjective assay scales of pain including, e.g., the Pain Intensity Visual Analogue Scale or Pain Intensity Categorical Scale. Likewise, levels of “pain relief” can also be quantified by a subjective assay, e.g., Time to 10 Perceptible and Meaningful Pain Relief.

The terms “intravenous formulation,” or “intravenous acetaminophen formulation” shall refer to a single dose formulation of acetaminophen that is provided as a lyophilized powder (or other solid form) that, once reconstituted in solution, is physiologically compatible with intravenous administration (e.g., by injection, infusion or otherwise). Alternatively, the terms refer to a formulation that is provided as a solution.

Reduced Dose Acetaminophen Formulations for Intravenous Administration (IV Formulations)

In some embodiments, the IV acetaminophen formulations described herein are in the form of a lyophilized powder to be reconstituted in solution under sterile conditions prior to administration. In other embodiments, the IV acetaminophen formulations are provided as sterile solutions ready for administration. Appropriate containers (e.g., vials, bottles, ampules, containers, etc.) for the IV formulations in either of the forms just described, as well as aseptic techniques are well known.

IV Acetaminophen Dosage

In various embodiments, the single dose IV acetaminophen formulation contains less than about 1 gram of acetaminophen. In some embodiments, the single dose IV acetaminophen contains about 500 to about 1000 mgs. In some 35 embodiments, the single dose IV acetaminophen contains about 550 mgs to about 900 mgs. In some embodiments, the single dose IV acetaminophen formulations described herein contain about 550 mg to about 800 mg of acetaminophen, i.e., about 560 mg, 570 mg, 580 mg, 600 mg, 610 mg, 620 40 mg, 630 mg, 640 mg, 650 mg, 660 mg, 670 mg, 675 mg, 680 mg, 690 mg, 700 mg, 720 mg, 750 mg, 775 mg, or any other amount of acetaminophen from about 550 mg to about 800 mg of acetaminophen. In some embodiments, an IV acetaminophen formulation contains about 600 mg to about 700 45 mg of acetaminophen, i.e., about 610 mg, 620 mg, 630 mg, 640 mg, 650 mg, 660 mg, 670 mg, 680 mg, 690 mg, or any other amount of acetaminophen from about 600 mg to about 700 mg of acetaminophen. In one embodiment, the acetaminophen formulation contains about 650 mg of acetaminophen.

In some embodiments, the concentration of acetaminophen in an IV formulation solution described herein is about 0.3% (w/v) to about 12% (w/v), i.e., about 0.4%, 0.5%, 0.6%, 0.7%, 0.8%, 0.9%, 1.0%, 1.1%, 1.2%, 1.5%, 2.5%, 55 3%, 3.7%, 4%, 4.5%, 5%, 6%, 7%, 8%, 8.5%, 9%, 10%, 10.5%, 11%, or any other concentration from about 0.3% (w/v) to about 12% (w/v). In some embodiments the concentration of acetaminophen is about 0.7% (w/v) to about 1.4% (w/v), i.e., about 0.8%, 0.9%, 1.0%, 1.1%, 1.2%, 1.3% 60 or any other concentration of acetaminophen from about 0.7% (w/v) to about 1.4% (w/v). In one embodiment, the concentration of acetaminophen is about 1.0% (w/v).

In some embodiments, the volume of an IV acetaminophen formulation solution is about 30 to about 200 ml, i.e., about 30, 35, 40, 45, 55, 60, 65, 75, 80, 85, 90, 92, 95, 100, 105, 110, 125, 130, 150, 175, 180, or another volume of IV 65 formulation solution from about 30 to about 200 ml. In some

US 9,610,265 B2

7

embodiments, the volume of the IV formulation is about 75 to about 125 ml. In another embodiment the volume is about 40 to about 75 ml. In one embodiment, the volume of the IV formulation is about 100 ml.

Antioxidants

Generally, the acetaminophen formulations described herein also contains at least one antioxidant to increase the stability of acetaminophen in solution. Examples of suitable antioxidants include, but are not limited to, cysteine hydrochloride monohydrate, thioglycolic acid, thiolacetic acid, dithiothreitol, reduced glutathione, thiourea, alpha-thioglycerol, cysteine, acetylcysteine, methionine, mercaptoethane sulfonic acid, metabisulfite, ascorbic acid ascorbic acid derivatives (e.g., ascorbyl palmitate), sodium citrate, an organic compound having at least one thiol, an alkyl polyhydroxylated compound, a cycloalkyl polyhydroxylated compound, a hydroxypolycarboxylic acid, an alpha-hydroxypolycarboxylic acid (e.g., citric acid), tocotrienol, dimethyl glycine, betaine, butylated hydroxyanisole, butylated hydroxytoluene, tocopherol, tocopherol polyethylene glycol succinate, butylated hydroxytoluene (BHT), butylated hydroxyanisole (BHA), propyl gallate, hydroquinone, hydroxycoumarins, ethanolamine, lecithin, cephalin, malic acid, sorbitol, phosphoric acid, thiopropionic acid and its esters, dithiocarbamates or any combination thereof. In one embodiment, the acetaminophen formulation is free of polyethylene glycol or a derivative thereof. In another embodiment, the acetaminophen formulation is free of sulfites. In one embodiment, the antioxidant is cysteine hydrochloride monohydrate. In yet another embodiment, the antioxidant is mannitol.

In some embodiments, the amount % (w/w) of the antioxidant in the solid form of the IV formulation (i.e., prior to preparation in solution) is about 0.10% (w/w) to about 5.0% (w/w), i.e., 0.15% (w/w), 0.17% (w/w), 0.20% (w/w), 0.30% (w/w), 0.40% (w/w), 0.45% (w/w), 0.50% (w/w), 0.52% (w/w), 0.55% (w/w), 0.60% (w/w), 0.70% (w/w), 0.80% (w/w), 1.0% (w/w), 1.3% (w/w), 1.5% (w/w), 1.7% (w/w), 2.0% (w/w), 2.2% (w/w), 2.3% (w/w), 2.5% (w/w), 2.7%, 2.8%, 3.0% (w/w), 3.2%, 3.5% (w/w), 3.6% (w/w), 4.0% (w/w), 4.7% (w/w), or any other amount of antioxidant % (w/w) from about 0.10% (w/w) to about 5.0% (w/w). In some embodiments, the amount % (w/w) of antioxidant is about 0.30% (w/w) to about 1.0% (w/w). In one embodiment, the amount % (w/w) of antioxidant is about 0.50% (w/w).

In some embodiments, the concentration of the antioxidant in an IV formulation solution prior to administration ranges from about 0.01 mg/ml to about 10 mg/ml, i.e., 0.02 mg/ml, 0.03 mg/ml, 0.05 mg/ml, 0.08 mg/ml, 0.09 mg/ml, 0.10 mg/ml, 0.12 mg/ml, 0.13 mg/ml, 0.15 mg/ml, 0.18 mg/ml, 0.20 mg/ml, 0.22 mg/ml, 0.25 mg/ml, 0.27 mg/ml, 0.30 mg/ml, 0.40 mg/ml, 0.45 mg/ml, 0.50 mg/ml, 0.60 mg/ml, 0.80 mg/ml, 1.2 mg/ml, 1.5 mg/ml, 2.0 mg/ml, 2.5 mg/ml, 3.0 mg/ml, 3.5 mg/ml, 4.0 mg/ml, 5.0 mg/ml, 6.0, mg/ml 7.5 mg/ml, 8.0 mg/ml, 9 mg/ml, 9.5 mg/ml, or any other concentration of antioxidant from about 0.01 mg/ml to about 10 mg/ml. In some embodiments, the concentration of antioxidant is about 0.08 mg/ml to about 0.50 mg/ml. In one embodiment, the concentration of antioxidant is about 0.25 mg/ml.

Buffering Agents

In some embodiments, an IV acetaminophen formulation contains at least one buffering agent to maintain the pH of the formulation within an acceptable range as described herein. The buffer used is a buffer compatible with parenteral administration in humans, the pH of which may be adjusted

8

between 4 and 8. In some embodiments, the pH of an IV acetaminophen formulation is from about pH 4 to about pH 8, i.e., pH 4.5, pH 4.6, pH 4.8, pH 5.0, pH 5.5, pH 6.2, pH 6.5, pH 7.5, or any other pH value from about pH 4 to about

5 pH 8. In some embodiments, the pH of the IV acetaminophen formulation is from about pH 5 to about pH 7.0, i.e., about pH 5.2, pH 5.5, pH 5.6, pH 6.0, pH, 6.4, or any other pH value from about pH 5 to about pH 7.0. In one embodiment, the IV acetaminophen formulation has a pH of about 10 5 to about 6.

In some embodiments, buffering agents have a pKa from about 4.5 to about 6.5, i.e., 4.6, 4.8, 5.0, 5.2, 5.3, 5.4, 5.5, 15 5.8, 6.0, 6.2, 6.4, or any other pKa from about 4.5 to about 6.5.

In some embodiments, the buffering agent is a pharmaceutically acceptable salt or acid of citrate, phosphate, acetate, glutamate, tartrate, benzoate, lactate, histidine or other amino acids, gluconate, malate, succinate, formate, 20 propionate, carbonate, or any combination thereof adjusted to an appropriate pH, as described herein, with acid (e.g., hydrochloric acid) or base (e.g., sodium hydroxide) as required. In one embodiment, the buffering agent is disodium phosphate dehydrate.

25 In some embodiments, the amount % (w/w) of the buffering agent in the solid form of the IV formulation (i.e., prior to preparation in solution) is about 0.05% (w/w) to about 2% (w/w), i.e., about 0.08% (w/w), 0.10% (w/w), 0.15% (w/w), 1.0% (w/w), 1.3% (w/w), 1.5% (w/w), 1.7% (w/w), 0.20% 30 (w/w), 0.22% (w/w), 0.25% (w/w), 0.26% (w/w), 0.27% (w/w), 0.28% (w/w), 0.30% (w/w), 0.35% (w/w), 0.40% (w/w), 0.50% (w/w), 0.60% (w/w), 0.70% (w/w), 0.80% (w/w), 1.2% (w/w), 1.4% (w/w), 1.5% (w/w), 1.7%, or any other amount of buffering agent % (w/w) from about 0.05% 35 (w/w) to about 2.0% (w/w). In some embodiments, the amount % (w/w) of the buffering agent is about 0.10% to about 0.70%. In one embodiment, the amount % (w/w) of the buffering agent is about 0.26%.

40 In some embodiments, the concentration of the buffering agent in an IV formulation solution prior to administration ranges from about 0.01 mg/ml to about 10 mg/ml, i.e., 0.02 mg/ml, 0.03 mg/ml, 0.05 mg/ml, 0.08 mg/ml, 0.09 mg/ml, 0.10 mg/ml, 0.12 mg/ml, 0.13 mg/ml, 0.15 mg/ml, 0.30 mg/ml, 0.5 mg/ml, 0.8 mg/ml, 1.2 mg/ml, 1.5 mg/ml, 2.0 mg/ml, 2.5 mg/ml, 3.0 mg/ml, 3.5 mg/ml, 4.0 mg/ml, 5.0 mg/ml, 5.0 mg/ml, 6.0, mg/ml 7.5 mg/ml, 8.0 mg/ml, 9 mg/ml, 9.5 mg/ml, or any other concentration of buffering agent from about 0.01 mg/ml to about 10 mg/ml. In some embodiments, the concentration of buffering agent is about 0.08 mg/ml to about 0.30 mg/ml. In one embodiment, the concentration of buffering agent is about 0.13 mg/ml.

Isotonicity Agents

50 In some embodiments, an IV acetaminophen formulation also contains one or more isotonicity agents to maintain the 55 osmolality of the formulation in a range that is physiologically compatible with IV administration. In some embodiments, the osmolality of the IV acetaminophen formulation is about 230 mOsm/L to about 420 mOsm/L, i.e., about 240 mOsm/L, 250 mOsm/L, 260 mOsm/L, 270 mOsm/L, 280 mOsm/L, 290 mOsm/L, 300 mOsm/L, 305 mOsm/L, 310 mOsm/L, 320 mOsm/L, 350 mOsm/L, 375 mOsm/L, 400 mOsm/L or any other osmolality from about 240 mOsm/L to about 420 mOsm/L. In some embodiments, the osmolality of the IV acetaminophen formulation is about 280 mOsm/L to about 320 mOsm/L, i.e., about 290 mOsm/L, 295 mOsm/L, 300 mOsm/L, 305 mOsm/L, 310 mOsm/L, 315 mOsm/L, or any other osmolality from about 280 mOsm/L to about 320

US 9,610,265 B2

9

mOsm/L. In one embodiment, the osmolality of the IV acetaminophen formulation is about 200-400 mOsm/L.

Suitable agents for adjusting the isotonicity of IV acetaminophen formulations include, but are not limited to, mannitol, sorbitol, glycerol, sucrose, glucose, dextrose, levulose, fructose, lactose, polyethylene glycols 400 to 4000, phosphates, sodium chloride, potassium chloride, calcium chloride, calcium gluconoglucoheptonate, dimethyl sulfone. In one embodiment, the isotonicity agent is mannitol.

In some embodiments, the amount % (w/w) of the isotonicity agent in the solid form of the IV formulation (i.e., prior to preparation in solution) is about 5% (w/w) to about 95% (w/w), i.e., about 10% (w/w), 15% (w/w), 20% (w/w), 25% (w/w), 30% (w/w), 35% (w/w), 40% (w/w), 45% (w/w), 50% (w/w), 55% (w/w), 60% (w/w), 65% (w/w), 70% (w/w), 72% (w/w), 74% (w/w), 76% (w/w), 78% (w/w), 79% (w/w), 80% (w/w), 81% (w/w), 82% (w/w), 84% (w/w), 86% (w/w), 90% (w/w), 92% (w/w), or any other amount of isotonicity agent % (w/w) from about 5% (w/w) to about 95% (w/w). In some embodiments, the amount of isotonicity agent % (w/w) is about 65% (w/w) to about 85% (w/w). In one embodiment, the amount of isotonicity agent % (w/w) is about 79%.

In some embodiments, the concentration of the isotonicity agent in an IV formulation solution prior to administration ranges from about 1.0 mg/ml to about 150 mg/ml, i.e., 1.0 mg/ml, 2.0 mg/ml, 3.0 mg/ml, 3.5 mg/ml, 4.0 mg/ml, 4.5 mg/ml, 5.0 mg/ml, 8.0 mg/ml, 12 mg/ml, 15 mg/ml, 20 mg/ml, 25 mg/ml, 30 mg/ml, 32 mg/ml, 35 mg/ml, 37 mg/ml, 38 mg/ml, 40 mg/ml, 50 mg/ml, 60 mg/ml, 75 mg/ml, 80 mg/ml, 90 mg/ml, 95 mg/ml, 100, 110, 120, 140, or any other concentration of buffering agent from about 5 mg/ml to about 150 mg/ml. In some embodiments, the concentration of buffering agent is about 0.08 mg/ml to about 0.30 mg/ml. In one embodiment, the concentration of buffering agent is about 0.13 mg/ml.

Stabilizers

In some embodiments, IV acetaminophen formulations described herein also include a stabilizer, e.g., a chelating agent such as ethylene diamino tetraacetic acid (EDTA), ethylene diamino, N,N'-diacetic-N,N'-dipropionic acid, ethylene diamino tetraphosphonic acid, 2,2'-(ethylene diamino) dibutyric acid, nitrilotriacetic acid, or ethylene-glycol bis (diaminoethyl ether) N,N,N',N'-tetraacetic acid and sodium or calcium salts thereof. In some embodiments, the IV acetaminophen formulation includes EDTA as the stabilizer.

In some embodiments, the IV acetaminophen formulations described herein contain a stabilizer in the amount of about 0.005 to about 1.0 mg/ml. In some embodiments, the stabilizer is present in an amount of about 0.01 mg/ml, 0.05 mg/ml, 0.1 mg/ml, 0.5 mg/ml, or 1.0 mg/ml.

In some embodiments, to reduce oxidation of acetaminophen in solution and thereby increase its stability, oxygen is removed from an IV formulation solution by bubbling an inert gas (e.g., argon or nitrogen) through the solution under sterile conditions. Methods for minimizing oxidative degradation of acetaminophen solutions during storage are described in further detail in, e.g., U.S. Pat. No. 6,992,218, which is incorporated herein by reference in its entirety.

Methods of Treatment

In many cases, IV administration of acetaminophen is considered the most suitable route of administration for expedient and efficacious relief of a patient's pain or fever, particularly in a hospital setting. In some embodiments, a subject to be administered an IV formulation of acetaminophen (e.g., an adult subject or adolescent weighing at least

10

about 50 kg), as described herein, is unconscious, sedated, fasting, nauseous, or unable to be administered a pharmaceutical composition by an oral route. Additionally, the rectal route is associated with highly variable bioavailability and slow absorption, and in children, the efficacious rectal dose exposes some pediatric patients to a potentially toxic exposure. In some embodiments, a patient suffering from pain or fever is in need of a faster onset of pain relief or fever treatment than possible by acetaminophen administration through an administration route other than by an IV administration.

In some embodiments, the IV formulations described herein are used as a pretreatment to another therapy. In some of these embodiments, pretreatment with an IV formulation described herein allows the use of a lower dose of acetaminophen. In some embodiments, the IV formulation described herein is administered before chemotherapy treatment, radiation treatment, a biopsy, or a blood transfusion. It should be understood that these are non-limiting examples and that the IV formulations described herein can be administered as a pretreatment to any therapy where pain and/or fever are predicted to occur.

The IV formulations described herein can be used for reducing pain conditions including, but not limited to, acute nociceptive pain, acute neuropathic pain, postoperative pain, lower back pain, cluster headaches, herpes neuralgia, phantom limb pain, central pain, dental pain, opioid-resistant pain, visceral pain, surgical pain, procedural pain, bone injury pain, pain during labor and delivery, pain resulting from burns, post partum pain, headache, muscular aches, backache, arthritis pain, the common cold, toothache, dental pain, osteoarthritis pain, menstrual pain, menstrual cramps, migraine, and genitourinary tract-related pain including cystitis. In some embodiments, the IV formulation is administered preemptively to a subject, i.e., prior to the onset of pain or a pain-inducing condition or stimulus (e.g., a surgical operation). In some embodiments, the IV formulations described herein are used to reduce fever, including, but not limited to, fever due to infections, drug reactions, allergic reactions, transfusion reactions, stroke, surgery, heat stroke, rheumatic diseases, cancer, or fever of unknown origin. In some embodiments, the IV formulations described herein are administered to a patient undergoing a dental procedure.

In some embodiments, the IV formulation is administered to a subject after undergoing a surgical intervention, e.g., within about 12 hours after a surgical intervention, i.e., within 11 hours, 10 hours, 9 hours, 8 hours, 6 hours, 5 hours, 4 hours, 3 hours, 2 hours, 1 hours, 45 minutes, 30 minutes 15 minutes, 5 minutes, or any period within about 12 hours following a surgical intervention.

In some embodiments, a subject is administered the IV formulation prior to a surgical intervention, e.g., about 4 hours or less prior to the surgical intervention, i.e., about 3 hours, 2 hours, 1 hours, 30 minutes, 15 minutes or even during the surgical intervention itself.

Depending on the concentration of acetaminophen in an IV formulation solution and consistent with the acetaminophen dose levels described herein, the volume of IV formulation solution to be administered can vary from about 1 mL to about 200 mL, e.g., 5 mL, 10 mL, 20 mL, 25 mL, 30 mL, 40 mL, 50 mL, 60 mL, 65 mL, 70 mL, 85 mL, 90 mL, 100 mL, 110 mL, 120 mL, 130 mL, 140 mL, 150 mL, 160 mL, 180 mL, or any other volume of IV formulation solution from about 1 mL to about 200 mL.

In some embodiments, the amount of time required for administration of the IV formulation ranges from about 1 minute to about 1 hours, i.e., about 5 minutes, 10 minutes,

US 9,610,265 B2

11

11 minutes, 15 minutes, 20 minutes, 30 minutes, 45 minutes, or any other administration time from about 1 minute to about 1 hour. In some embodiments, the amount of time required for administration of the IV formulation ranges from about 5 minutes to about 45 minutes, or about 5 minutes to about 30 minutes, or about 5 minutes to about 15 minutes.

Depending on the severity and persistence of a subject's condition, and in accordance with a medical caregiver's judgment, an IV formulation dose of acetaminophen, as described herein, can be administered in an interval to allow for the administration of about 3 to about 5 grams in a 24 hour period. In some embodiments, the IV formulation of acetaminophen is administered in an interval sufficient to allow for the administration of about 4 grams in a 24 hour period. In some embodiments, the IV formulation is administered between 1 to 6 times, i.e., 1, 2, 3, 4, 5, 6 times every twenty four hours, as deemed necessary by a medical caregiver. In some embodiments, the frequency of administration is not greater than once every four hours.

In various embodiments, the IV formulation of acetaminophen is dosed so as to provide less than about 4 grams over a 24 hour period. In various embodiments, the IV formulation of acetaminophen is dosed three to six times in a 24 hour period. For example, in some embodiments, the IV formulation of acetaminophen is dosed three times in a 24 hour period. In other embodiments, the IV formulation of acetaminophen is dosed four times in a 24 hour period. In still other embodiments, the IV formulation of acetaminophen is dosed five times in a 24 hour period. In some embodiments, the IV formulation of acetaminophen is dosed six times in a 24 hour period. In some embodiments, the IV formulation of acetaminophen is dosed seven times in a 24 hour period. In some embodiments, the IV formulation of acetaminophen is dosed eight times in a 24 hour period.

Combination Therapies

The acetaminophen IV formulations described herein can also be used in combination with other therapeutic reagents, e.g., other analgesics, antipyretics, or anti-inflammatory agents that are selected for their therapeutic or palliative value. In general, where a combination therapy is employed, other agents do not have to be administered in the same pharmaceutical composition as acetaminophen, and may, because of different physical and chemical characteristics, be administered by different routes. The determination of the mode of administration and the advisability of administration, where possible, in the same pharmaceutical composition, is well within the knowledge of the skilled clinician with the teachings described herein. The initial administration of either the IV acetaminophen formulation or the one or more therapeutic agents (e.g., analgesic agents other than acetaminophen) to be used in combination with acetaminophen can be made according to established protocols known in the art, and then, based upon the observed effects, the dosage, modes of administration and times of administration can be modified by the skilled clinician.

The particular choice of compounds (e.g., analgesic agents) for use in combination with the IV acetaminophen formulation described herein will depend on the diagnosis of the attending physicians (or other medical caregivers) and their judgment of the condition of the patient and the appropriate treatment protocol. The compounds may be administered concurrently (e.g., simultaneously, essentially simultaneously or within the same treatment protocol) or sequentially, depending upon the severity of pain experienced by the patient, the nature of the disease, disorder, or condition, the condition, and the actual choice of compounds

12

used. The determination of the order of administration, and the number of repetitions of administration of each therapeutic agent during a treatment protocol, is well within the knowledge of the skilled physician after evaluation of the disease being treated and the condition of the patient.

For combination therapies described herein, dosages of the compounds to be co-administered with an acetaminophen IV formulation will vary depending on the type of co-drug employed, on the amount of pain experienced by the patient, the risk for addiction, the disease or condition being treated and so forth. In addition, when co-administered with one or more biologically active agents, the acetaminophen IV formulation provided herein may be administered either simultaneously with the biologically active agent(s), or sequentially. If administered sequentially, the attending physician will decide on the appropriate sequence of administering protein in combination with the biologically active agent(s).

In any case, the multiple therapeutic agents (one of which is an acetaminophen IV formulation described herein) may be administered in any order or even simultaneously. If simultaneously, the multiple therapeutic agents may be provided in a single, unified IV form, or in multiple forms (by way of example only, either as a single IV formulation, as multiple IV formulations, or as IV formulation and a pill). One of the therapeutic agents may be given in multiple doses, or both may be given as multiple doses. If not simultaneous, the timing between the multiple doses may vary from more than 1 minute to less than 12 hours. In some embodiments, the timing between the multiple doses is from between about 1 minute to about 6 hours, or about 1 minute and about 3 hours, or about 1 minute and about 1 hour. In addition, the combination methods, compositions and formulations are not to be limited to the use of only two agents; the use of multiple therapeutic combinations is also envisioned.

The pharmaceutical agents which make up the combination therapy disclosed herein may be a combined dosage form (i.e., a combined IV formulation) or in separate dosage forms intended for substantially simultaneous administration. The pharmaceutical agents that make up the combination therapy may also be administered sequentially, with either therapeutic compound being administered by a regimen calling for two-step administration. The two-step administration regimen may call for sequential administration of the active agents or spaced-apart administration of the separate active agents. The time period between the multiple administration steps may range from, a few minutes to several hours, depending upon the properties of each pharmaceutical agent, such as potency, solubility, bioavailability, plasma half-life and kinetic profile of the pharmaceutical agent.

The compounds described herein and combination therapies can be administered before, during or after the occurrence of a fever or painful condition, and the timing of administering the composition containing a compound can vary. Thus, for example, the compounds can be used as a prophylactic and can be administered continuously to subjects with a propensity to develop conditions (e.g., body aches and chills following chemotherapy treatment) or diseases in order to prevent the occurrence of the disease or condition. The compounds and compositions can be administered to a subject during or as soon as possible after the onset of the symptoms. The administration of the compounds can be initiated within the first 48 hours of the onset of the symptoms, preferably within the first 48 hours of the onset of the symptoms, more preferably within the first 6

US 9,610,265 B2

13

hours of the onset of the symptoms, and most preferably within 3 hours of the onset of the symptoms.

A compound is preferably administered as soon as is practicable before or after the onset of a painful condition (e.g., postoperative pain), and for a length of time necessary for the treatment of the disease, such as, for example, from about 1 month to about 3 months.

Exemplary Analgesic Agents for Use in Combination with an Acetaminophen IV Formulation

Opioids

In some embodiments, an acetaminophen IV formulation described herein is used in any combination with one or more opioids, which include, but are not limited to allylpromidine, alphamethylfentanyl, alfentanil, bezitramide, buprenorphine, butorphanol, carfentanyl, codeine, dextropropoxyphene, dextromoramide, dezocine, diacetylmorphine, dihydrocodeine, dipapanone, morphine, dihydrocodeine, etorphine, fentanyl, hydrocodone, hydromorphone, ketobemidone, lefetamine, levorphanol, levo-alphacetylmethadol, levomethorphan, meptazinol, methadone, morphine, nalbuphine, nicomorphine, ohmefentanyl, opium, oripavine, oxycodone, oxymorphone, methadone, PEPAP, pentazocine, pethidine, phenazocine, piritamide, prodine, propoxyphene napsylate, remifentanil, sufentanil, tilidine, thebaine, tramadol, and tapentadol.

Non-Steroidal Anti-Inflammatory Drugs (NSAIDs)

In some embodiments, an acetaminophen IV formulation described herein is used in any combination with one or more NSAIDs, which include, but are not limited to amoxiprin, benorilate, choline magnesium salicylate, diflusinal, faislamine, methyl salicylate, magnesium salicylate, salicyl salicylate, diclofenac, aceclofenac, acemetacin, bromfenac, ethenzamide, etodolac, indometacin, nabumetone, sulindac, tolmetin, carprofen, fenbufen, fenoprofen, flurbiprofen, ketoprofen, ketorolac, loxoprofen, naproxen, oxaprozin, tiaprofenic acid, suprofen, mefenamic acid, meclofenamic acid, phenylbutazone, metamizole, oxyphenbutazone, sulfinpyrazone, piroxicam, lornoxicam, meloxicam, tenoxicam, nimesulide salixylates, arylalkanoic acids, 2-arylpropanoic acids (profens), n-arylanthranilic acids (fenamic acids), pyrazolidine derivatives, oxicams, and COX-2 inhibitors.

Other Analgesic Agents

In some embodiments, an acetaminophen IV formulation described herein is used in any combination with one or more analgesic agents not described above, including, but not limited to, barbiturates (e.g., butalbital), pyrazolones (e.g., aminophenazone, metamizole, phenazone), cannabinoids (e.g., tetrahydrocannabinol), ziconotide, choline magnesium fentanyl, oxycodone, codeine, dihydrocodeine, dihydrocodeineone enol acetate, morphine, desomorphine, apomorphine, diamorphine, pethidine, methadone, dextropropoxyphene, pentazocine, dextromoramide, oxymorphone, hydromorphone, dihydromorphone, noscapine, papaverine, papaveretum, alfentanil, buprenorphine, tramadol and pharmaceutically acceptable salts, derivatives, homologs or analogs thereof as well as opioid agonists. Exemplary Antiemetic Agents for Use in Combination with an Acetaminophen IV Formulation

In some embodiments, an acetaminophen IV formulation described herein is used in any combination with one or more antiemetic agents not described above, including, but not limited to, antihistamines (e.g., Cyclizine, Diphenhydramine, Dimenhydrinate, Meclizine, Promethazine, Pentazine, Phenergan, Promacot, or Hydroxyzine); 5-HT₃ receptor antagonists (e.g., Dolasetron, Granisetron, Ondansetron, Tropisetron, or Palonosetron); and dopamine

14

antagonists (e.g., Domperidone, Droperidol, haloperidol, chlorpromazine, promethazine, prochlorperazine, or metoclopramide).

Kits

In some embodiments provided herein are kits that can simplify the administration of an IV acetaminophen formulation to a patient. In some embodiments, a kit comprises a unit dosage form of an acetaminophen IV formulation as described herein provided as a sterile lyophilate to be reconstituted by addition of sterile water. In other embodiments, the IV formulation is provided as a sterile degassed solution ready for administration. The kit can further comprise a label or printed instructions on the use of the acetaminophen IV formulation to treat pain or fever. The kit can also further comprise a unit dosage form of another therapeutic agent, for example, a container containing an effective amount of a second analgesic agent for use in combination with the acetaminophen IV formulation. In some embodiments, a kit further comprises a device that is useful for administering the IV formulation unit dosage forms. Examples of such a device include, but are not limited to, a syringe or a drip bag.

While preferred embodiments of the present invention have been shown and described herein, it will be understood that such embodiments are provided by way of example only. Numerous variations, changes, and substitutions can be made without departing from the scope of the invention. It should be understood that various alternatives to the embodiments of the invention described herein may be employed in practicing the invention. It is intended that the following claims define the scope of the invention and that methods and structures within the scope of these claims and their equivalents be covered thereby. Thus, these examples should not be read as limiting the example in any way. For example, different amounts of the components described in the following examples as well as the components themselves can be changed according to the disclosure provided herein.

EXAMPLES

Example 1

IV Acetaminophen Formulations

TABLE 1

Exemplary IV Formulation of Acetaminophen		
Acetaminophen	0.550 g-1.000 g	
Excipients:		
Antioxidant	0.0100-0.0200 g	
pH modulator(s)	qs pH 5-6	
Buffer	0.005-0.01 g	
Isotonic Agent	1.5-3.5 g	
Solvent	qs 50.0-100.0 mL	

Example 1A

IV Acetaminophen Formulations

Example 1A is prepared according to the procedure outlined in Example 2 using the amounts of the excipients described in Example 1.

US 9,610,265 B2

15

16

Formula 1(A)					
	Acetaminophen				
Excipients:	0.550 g	0.650 g	0.750 g	0.850 g	0.950 g
Antioxidant	Reduced Glutathione				
pH Modulator	Sodium hydroxide				
pH Modulator	Hydrochloric Acid				
Buffering Agent	Sodium Citrate				
Isotonicity Agent	Sodium Chloride				
Solvent	Sterile Water for injection				

Example 1B

20

IV Acetaminophen Formulations

Example 1B is prepared according to the procedure outlined in Example 2 using the amounts of the excipients described in Example 1.

Formula 1(B)					
	Acetaminophen				
Excipients:	0.550 g	0.650 g	0.750 g	0.850 g	0.950 g
Antioxidant	Methionine	Methionine	Methionine	Methionine	Methionine
pH Modulator	Sodium hydroxide				
pH Modulator	Hydrochloric Acid				
Buffering Agent	Sodium Acetate				
Isotonicity Agent	Mannitol	Mannitol	Mannitol	Mannitol	Mannitol
Solvent	Sterile Water for injection				

Example 1C

IV Acetaminophen Formulations

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Example 1C is prepared according to the procedure outlined in Example 2 using the amounts of the excipients described in Example 1.

Formula 1(C)					
	Acetaminophen				
Excipients:	0.550 g	0.650 g	0.750 g	0.850 g	0.950 g
Antioxidant	Cysteine Hydrochloride				
pH Modulator	Sodium Monohydrate				
pH Modulator	Hydrochloric Acid				
Buffering Agent	Disodium Phosphate				
Isotonicity Agent	Mannitol	Mannitol	Mannitol	Mannitol	Mannitol
Solvent	Sterile Water for injection				

US 9,610,265 B2

17
Example 1D

18

IV Acetaminophen Formulations

Example 1D is prepared according to the procedure outlined in Example 2 using the amounts of the excipients described in Example 1. 5

Excipients:	Formula 1(D)				
	Acetaminophen				
	0.550 g	0.650 g	0.750 g	0.850 g	0.950 g
Antioxidant	Ascorbic Acid				
pH Modulator	Sodium hydroxide				
pH Modulator	Hydrochloric Acid				
Buffering Agent	Sodium Tartate				
Isotonicity Agent	Glycerol	Glycerol	Glycerol	Glycerol	Glycerol
Solvent	Sterile Water for injection				

Example 1E

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IV Acetaminophen Formulations

Example 1E is prepared according to the procedure outlined in Example 2 using the amounts of the excipients described in Example 1.

Excipients:	Formula 1(E)				
	Acetaminophen				
	0.550 g	0.650 g	0.750 g	0.850 g	0.950 g
Antioxidant	Acetylcysteine	Acetylcysteine	Acetylcysteine	Acetylcysteine	Acetylcysteine
pH Modulator	Sodium hydroxide				
pH Modulator	Hydrochloric Acid				
Buffering Agent	Disodium Phosphate				
	Dehydrate	Dehydrate	Dehydrate	Dehydrate	Dehydrate
Isotonicity Agent	Sorbitol	Sorbitol	Sorbitol	Sorbitol	Sorbitol
Solvent	Sterile Water for injection				

Example 1F

50

IV Acetaminophen Formulations

Example 1F is prepared according to the procedure outlined in Example 2 using the amounts of the excipients described in Example 1.

Excipients:	Formula 1(F)				
	Acetaminophen				
	0.550 g	0.650 g	0.750 g	0.850 g	0.950 g
Antioxidant	metabisulfite	metabisulfite	metabisulfite	metabisulfite	metabisulfite
pH Modulator	Sodium hydroxide				
pH Modulator	Hydrochloric Acid				

Formula 1(F)					
Acetaminophen					
Excipients:	0.550 g	0.650 g	0.750 g	0.850 g	0.950 g
Buffering Agent	Disodium Phosphate				
Isotonicity Agent	Dehydrate Glucose				
Solvent	Sterile Water for injection				

Example 2

Preparation of IV Formulation Solutions

Prior to storage the formulations set forth in Example 1 are subjected to bubbling with nitrogen, transferred to Type II colorless bottles, and then placed under vacuum (low pressure approx. 550 mm of Hg) before stoppering the bottles with a synthetic elastomer grey stopper crimped with an aluminum cap. The residual oxygen content is approximately 1.5 ppm of dissolved oxygen. The bottles are then sterilized at 121° C. for 15 minutes. Sterile solutions are stored at ambient temperature (less than 30° C.) for up to two years prior to use.

Example 3

A Phase III Randomized, Double-Blind, Placebo-Controlled, Multi-Center, Parallel-Group, Repeated-Dose Study of the Analgesic Efficacy and Safety of 650 mg IV Acetaminophen Versus Placebo for the Treatment of Postoperative Pain After Abdominal Laparoscopic Surgery

In an effort to provide an intravenous, non-NSAID, non-opioid treatment for pain relief, the safety and efficacy of a 650 mg IV dose of APAP for the treatment of acute pain is examined.

Study Design and Evaluation

A Phase III, randomized, double-blind, Placebo-controlled, multi-center, parallel-group, repeated dose study is conducted in approximately 240 Subjects who have undergone planned or elective abdominal laparoscopic surgery. Approximately 15 to 20 US sites will participate in the Study.

Subjects will be centrally randomized, across all study centers, to receive infusions of Study Medication (either APAP or Placebo) at a dose at a dose (650 mg, 1000 mg, or placebo) and schedule described below.

Timed PI and pain relief (PR) Assessments will begin at baseline just prior to T0, the start of the first infusion of Study Medication, and continue through T24 hours.

All Subjects have access to rescue medication at all times throughout the study, as described below.

The Study will include the following assessment periods and procedures:

Screening (Day-21 to Randomization)

Screening is the period that begins when the Subject signs the Informed Consent Form and ends with randomization to

15 Study Medication on POD1. During this period, the eligibility and baseline status of the Subject are determined.

Treatment Period (Dose 1/T0/POD1 to T24/POD2)

Administration of Study Medication (and Study-related assessments) will occur from T0 (morning of POD1) to T24 hours (morning of POD2).

Criteria for Evaluation

The primary efficacy endpoint is SPID24 (defined as the Sum of VAS score differences from baseline at T0 to T24), excluding all data after rescue medication.

Subject Selection Criteria

To be eligible for entry into the Study, Subjects must meet all of the following criteria prior to surgery: (1) Provide written Informed Consent prior to participation in the Study;

20 (2) is scheduled to undergo abdominal laparoscopic surgery (laparoscopic gastric bypass procedures are not eligible); (3)

If Subject is a female of childbearing potential, have a negative pregnancy test within 21 days of surgery; (4) be at least 18, but not more than 80 years of age; (5) Have a Body

35 Mass Index (BMI)≥19 and ≤45 lb/in²; (6) Have an ASA risk class of I, II, or III according to the American Society of Anesthesiologists; (7) Have the ability to read and understand the Study procedures and the use of the pain scales and have the ability to communicate meaningfully with the

40 Study Investigator and staff; (8) Be free of other physical, mental, or medical conditions which, in the opinion of the Investigator, makes Study participation inadvisable

Exclusion Criteria (Screening)

A Subject is NOT eligible for entry if ANY of the

45 following criteria are met: (1) Used opioids or tramadol daily for greater than 7 days prior to Study Medication administration (Subjects who, in the Investigator's opinion have or are developing opioid tolerance are to be excluded); (2) Has been treated with Chapparal, Comfrey, Germaner,

50 Gin Bu Huan, Kava, Pennyroyal, Skullcap, St. John's Wort, or Valerian within 14 days prior to surgery; (3) Has significant medical disease(s), laboratory abnormalities or condition(s) that in the Investigator's judgment could compromise the Subject's welfare, ability to communicate with the Study

55 staff, complete Study activities, or would otherwise contraindicate Study participation; (4) Has known hypersensitivity to opioids, acetaminophen, or the inactive ingredients (excipients) of the Study Medication; (5) Has known or suspected history of alcohol or drug abuse or dependence

60 within the previous 2 years; (6) Has impaired liver function, e.g., AST/ALT/bilirubin greater than or equal to 3.0 times the upper limit of normal, active hepatic disease, evidence of clinically significant liver disease, or other condition (e.g., alcoholism, cirrhosis, or hepatitis) that may suggest the

65 potential for an increased susceptibility to hepatic toxicity with Study Medication exposure; (7) Has been treated with monoamine oxidase inhibitors (MAOIs) within 7 days prior

US 9,610,265 B2

21

to surgery; (8) Has participated in another clinical Study (investigational or marketed product) within 30 days of surgery

Post Operative Exclusion Criteria

The Subject must not meet any of the following criteria after surgery and prior to randomization to Study Medication: (1) Had any other surgery than the planned laparoscopic surgery or had intra operative or post operative complications which in the view of the Investigator would make Study participation inadvisable; (2) Has taken non steroidal anti-inflammatory drugs (NSAIDs), steroids or MAOIs during the day after surgery. Exceptions: The use of low-dose aspirin, e.g., 81 mg/day, for cardioprophylaxis, and topical or inhaled steroids are acceptable; (3) Had any neuraxial opioids or continuous local anesthetic infusions via percutaneous catheters administered as part of the anesthetic or post operative analgesic management (local anesthetic infiltration of surgical wounds at the time of closure is acceptable if done as a single injection); (4) Had a fever (greater than 38.6° C. or 101.5° F.) requiring treatment.

Postoperative Assessment (POD0)

The Subject will undergo abdominal laparoscopic surgery or other approved surgical procedure as described herein. Details of the surgical procedure(s) will be recorded on the CRF including the type of procedure(s) performed and perioperative medication will be recorded.

Example 4

Phase III, Open-Label, Prospective, Multi-Center, Repeated Dose, Randomized, Multi-Day Safety and Efficacy Study of 650 mg IV Acetaminophen

A Phase III, open-label, prospective, multi-center, repeated dose, randomized, multi-day safety and efficacy study was conducted in 213 subjects. The subjects were randomized as follows: 92 subjects to a q6 group (1 g of IV acetaminophen every 6 hours), 91 subjects to a q4 group (650 mg of IV acetaminophen every 4 hours), and 28 subjects to a standard of care control group, which could include oral acetaminophen, but no IV acetaminophen. Subjects who completed 5 days of study treatment included 63 in the q6 group, 59 in the q4 group and 26 in the control group. The primary endpoint was an assessment of safety using spontaneous adverse event reporting and daily liver enzymes. Efficacy evaluations were also performed.

Inclusion Criteria (Screening)

To be eligible for entry into the Study, Subjects had to meet all the following criteria: (1) Provide written informed consent prior to participation in the Study; (2) Be at least 18 years of age and weigh at least 41 kg; (3) Be anticipated by the Investigator to require multi-day (target is five days) use of IV treatment either because of: (a) having a "nothing by mouth" (NPO) status, (b) having a medical condition that makes oral intake difficult, or (c) having a medical condition that requires IV treatment; (4) Be willing to undergo 5 days of treatment with IV acetaminophen for the treatment of pain or fever (defined as a core temperature $\geq 38^{\circ}$ C.). Subjects had a slightly less than 15% chance (one in seven) of being assigned to the Control Group and receiving standard of care treatment, but no IV APAP; (5) Have the ability to read and understand the Study procedures and have the ability to communicate meaningfully with the Study Investigator and staff; and (6) If a female of child bearing potential, have a negative pregnancy test within 48 hours of randomization.

22

Exclusion Criteria (Screening)

A Subject was not eligible for entry if any of the following criteria were met: (1) Had a significant medical disease, laboratory abnormality or condition that, in the Investigator's judgment, could compromise the Subject's welfare or would otherwise contraindicate Study participation; (2) Was expected to have difficulty in communicating with the Study staff or completing Study requirements (including follow up visits); (3) Had known hypersensitivity to acetaminophen or the inactive ingredients (excipients) of IV acetaminophen or any contraindication to receiving acetaminophen; (4) Had impaired liver function, e.g., ALT greater than or equal to 3 times the upper limit of normal (ULN), bilirubin greater than or equal to 3 times ULN, known active hepatic disease (e.g., hepatitis), evidence of clinically significant chronic liver disease or other condition affecting the liver (e.g., alcoholism as defined by DSM-IV, cirrhosis or chronic hepatitis); or (5) Had participated in an interventional clinical Study (investigational or marketed product) within 30 days of Study entry.

Efficacy Analysis

All analyses of efficacy were conducted on the modified intent-to-treat population separately for the two indications (acute pain and fever). Subjects' Global Evaluations were summarized descriptively (n, mean, SD, median, minimum, and maximum) by treatment group for each study day and for overall assessments. Summary statistics were also provided for each site.

Comparisons of efficacy endpoints between the following pairs of treatment groups were investigated using two-sided tests at the 5% level of significance:

IV acetaminophen 1 versus IV acetaminophen 650 mg
IV acetaminophen 1 g versus standard of care treatment
IV acetaminophen 650 mg versus standard of care treatment

A one-way analysis of variance (ANOVA) model with treatment group as the factor was used to test the treatment difference between these pairs. All groups were included in this analysis model. The p-values from the ANOVA model were presented along with the summary statistics.

Safety Analyses

All analyses of safety were conducted on the safety population.

Percentage of subjects withdrawn due to adverse event, percentage of subjects with adverse events (AEs) or serious adverse events (SAEs), and percentage of subjects with clinically meaningful changes in laboratory parameters were summarized.

All adverse events and serious adverse events were coded according to the Medical Dictionary for Regulatory Activities (MedDRA), Version 10.0. Additional analyses included displays of the number of subjects reporting at least one AE (incidence table), total number of episodes of each AE by body system and by severity, total number of episodes of each AE by body system, and by attribution. Liver function test abnormalities were graded using the Common Terminology Criteria for Adverse Events.

For each clinical laboratory parameter, descriptive statistics (n, mean, standard deviation, median, and range) were tabulated for baseline and final values. Change from baseline was tabulated for those subjects who had both baseline and final values. Liver function tests were also evaluated using values that were normalized to the upper limit of normal values for the local laboratory.

A shift table was prepared to present the shift in baseline clinical laboratory values that were clinically relevantly high or low at baseline and/or final measurement.

US 9,610,265 B2

23

Descriptive statistics (n, mean, standard deviation, median, and range) were tabulated for changes in vital signs from baseline to final measurement.

Results

Disposition of Subjects

A total of 257 subjects were screened for study enrollment. Of the total screened, 44 were screen failures, and 213 were enrolled and randomized: 92 subjects in the q6 group, 91 subjects in the q4 group, and 28 subjects in the control group. Subjects who completed 5 days of study treatment included 63 in the q6 group, 59 in the q4 group and 26 in the control group.

Subjects in the q4h group and q6h group were considered to be a Study Treatment Discontinuation/Early Termination if they received at least one dose of IV acetaminophen and discontinued study participation prior to completion of Day 5 treatments. Subjects in the control group were considered to be a Study Treatment Discontinuation/Early Termination if they discontinued at any time after T0, but prior to completion of Day 5 standard of care treatments.

Subjects who received at least one dose of IV acetaminophen and discontinued study participation prior to completing Day 5 treatments, but returned for the Last Study Visit were considered as "Partial Treatment Completers". Similarly, subjects in the control group who discontinued study participation prior to completing Day 5 standard of care treatments, but returned for the Last Study Visit were considered as "Partial Treatment Completers".

Subjects who completed Day 5 treatments (IV acetaminophen or standard of care) and procedures were characterized as a "Treatment Completer". A Treatment Completer who elected to discontinue study participation prior to the Last Study Visit was characterized as a Treatment Completer Early Termination.

Safety Outcome

There were no clinically relevant differences between the treatment groups in the frequency of serious, severe, related, or overall treatment emergent adverse events (TEAEs). In fact, most TEAEs were assessed by the Investigator to be mild or moderate in severity. The frequency of liver enzyme elevations seen in the treatment groups was comparable. More specifically, with regard to the hepatic transaminases alanine aminotransferase and aspartate aminotransferase, the frequency and severity of the elevations were comparable between the treatment groups. There were no clinically relevant differences between the treatment groups regarding laboratory assessments, vital signs, or physical examinations. Thus, based on these data, intravenous acetaminophen in both active treatment groups (i.e., 650 mg and 1000 mg dose groups) was well tolerated.

Efficacy Outcome

The modified intent-to-treat population was used for all analyses of efficacy: Subject Global Evaluations (rating of study treatments and rating of satisfaction with side effects related to study treatments) provided as a daily lookback (days 2 through 5) and overall evaluation (overall treatment period lookback) using a 4 point categorical rating scale (0=poor, 1=fair, 2=good, 3=excellent). A one-way ANOVA model with treatment group as the factor was used to test the treatment difference between each treatment pair:

IV acetaminophen 1 q6h versus IV acetaminophen 650 mg q4h

IV acetaminophen 1 g q6h versus standard of care treatment (Control)

IV acetaminophen 650 mg q4h versus standard of care treatment (Control)

24

All endpoints were tested at the 0.05 significance level (two-sided).

The IV acetaminophen 650 mg q4h group relative to the control group produced statistically significantly better satisfaction ratings for the Subject Global Assessments rating the level of satisfaction with the side effects related to study treatments the on the day 5 (mean rating 2.4 vs. 2.0, p=0.0167) and at the end of day 5 prior to discharge (mean rating 2.4 vs. 2.0, p=0.0129) 24 h look back assessments. On day 4, the satisfaction rating showed a trend to significance (mean rating 2.5 vs. 2.2, p=0.1162). With respect to the Subject Global Assessments rating the level of satisfaction with the study treatments, there were no statistically significant differences between the IV acetaminophen 650 mg q6h group and control group at any of the assessment points. For the both of the Subject Global Assessments rating either the level of satisfaction with the study treatments or the level of satisfaction with the side effects related to study treatments, there was no statistically significant differences between the two active treatment groups with respect to the daily 24 h lookback assessments on day 2, day 3, day 4, day 5, or at the end of day 5 prior to discharge; nor was there a statistically significant difference on the overall assessment at the Study Completion Visit.

The IV acetaminophen 1 g q6h group produced statistically significantly better satisfaction ratings for the Subject Global Assessments rating the level of satisfaction with the side effects related to study treatments on day 5 (mean rating 2.4 vs. 2.0, p=0.0062) and at the end of day 5 prior to discharge (mean rating 2.5 vs. 2.0, p=0.0073) 24 h lookback assessments compared to the control group. On day 4, the satisfaction rating showed a trend to significance (mean rating 2.5 vs. 2.2, p=0.0744). With respect to the Subject Global Assessments rating the level of satisfaction with the study treatments, there were no statistically significant differences between the IV acetaminophen 1 g q6h group and control group at any of the assessment points.

Statistically significant differences were observed for both active treatment groups versus the control group in the Subject Global Assessments rating the level of Satisfaction with the side effects related to study treatments on the day 5 and on the end of day 5 prior to discharge daily 24 h lookback assessments. Thus, these data suggest that the IV acetaminophen 1 q6h and 650 mg q4h groups were efficacious and provided comparable efficacy based upon the global satisfaction ratings.

Many modifications, equivalents, and variations of the present invention are possible in light of the above teachings, therefore, it is to be understood that within the scope of the appended claims, the invention may be practiced other than as specifically described.

What is claimed is:

1. A method of treating pain in a human subject weighing at least 50 kg comprising:

co-administering to the subject a therapeutically effective amount of a first pharmaceutical composition comprising about 500 mg to about 750 mg of acetaminophen and a therapeutically effective amount a second pharmaceutical composition comprising an opioid analgesic;

wherein the first pharmaceutical composition is administered to the subject intravenously.

2. The method of claim 1 wherein the administration of the first pharmaceutical composition to the subject is repeated at least once every four hours for a twenty-four hour period.

US 9,610,265 B2

25

3. The method of claim 1, wherein subject is administered less than 4 grams of acetaminophen over a twenty-four hour period.

4. The method of claim 3, wherein the first pharmaceutical composition is administered to the subject at least six times in a period of 24 hours.

5. The method of claim 3, wherein the first pharmaceutical composition is administered to the subject at least seven times in a period of 24 hours.

6. The method of claim 1, wherein the first pharmaceutical composition comprises about 600 mg to about 700 mg of acetaminophen.

7. The method of claim 1, wherein the first pharmaceutical composition comprises about 650 mg of acetaminophen.

8. The method of claim 1, wherein the opioid analgesic is selected from the group consisting of allylprodine, alphamethylfentanyl, alfentanil, bezitramide, buprenorphine, butorphanol, carfentanyl, codeine, dextropropoxyphene, dextromoramide, dezocine, diacetylmorphine, dihydrocodeine, dipapanone, dismorphine, dihydrocodeine, etorphine, fentanyl, hydrocodone, hydromorphone, ketobemidone, lefetamine, levorphanol, levo-alphaacetylmethodol, levomethorphan, meptazinol, methadone, morphine, nalbuphine, nicomorphine, ohmefentanyl, opium, oripavine, oxycodone, oxymorphone, methadone, phenethylphenylacetoxypiperidine (PÈPAP), pentazocine, pethidine, phenazocine, piritamide, prodine, propoxyphene napsylate, remifentanil, sufentanil, tilidine, thebaine, tramadol, and tapentadol.

9. The method of claim 8, wherein the opioid analgesic is selected from the group consisting of codeine, dihydrocodeine, hydrocodone, morphine, oxycodone, pentazocine, and tramadol.

10. The method of claim 8, wherein the opioid analgesic is morphine.

26

11. The method of claim 1, wherein the second pharmaceutical composition is administered to the subject orally, intravenously, rectally, or parenterally.

12. The method of claim 1, wherein the second pharmaceutical composition is administered as a single dose or in multiple doses.

13. The method of claim 1, wherein the first and second pharmaceutical compositions are administered simultaneously.

14. The method of claim 13, wherein the first and second pharmaceutical compositions are administered as a combined intravenous dosage form.

15. The method of claim 1, wherein the first and second pharmaceutical compositions are administered sequentially.

16. The method of claim 15, wherein the first pharmaceutical composition is administered prior to the second pharmaceutical composition or wherein the second pharmaceutical composition is administered prior to the first pharmaceutical composition.

17. The method of claim 15, wherein the timing between the administration of the first and second pharmaceutical compositions is from about 1 minute to less than about 12 hours.

18. The method of claim 15, wherein the timing between the administration of the first and second pharmaceutical compositions is from about 1 minute to less than about 6 hours.

19. The method of claim 15, wherein the timing between the administration of the first and second pharmaceutical compositions is from about 1 minute to less than about 3 hours.

20. The method of claim 15, wherein the timing between the administration of the first and second pharmaceutical compositions is from about 1 minute to less than about 1 hour.

* * * * *

EXHIBIT C

HIGHLIGHTS OF PRESCRIBING INFORMATION

These highlights do not include all the information needed to use OFIRMEV® safely and effectively. See full prescribing information for OFIRMEV.

OFIRMEV (acetaminophen) Injection

Initial U.S. Approval: 1951

WARNING: RISK OF MEDICATION ERRORS AND HEPATOTOXICITY

See full prescribing information for complete boxed warning

Take care when prescribing, preparing, and administering OFIRMEV Injection to avoid dosing errors which could result in accidental overdose and death.

OFIRMEV contains acetaminophen. Acetaminophen has been associated with cases of acute liver failure, at times resulting in liver transplant and death. Most of the cases of liver injury are associated with the use of acetaminophen at doses that exceed the recommended maximum daily limits, and often involve more than one acetaminophen-containing product (see WARNINGS).

RECENT MAJOR CHANGES

Boxed Warning	10/2013
Dosage and Administration	
General Dosing Information (2.1)	10/2013
Recommended Dosage: Adults and Adolescents (2.2)	10/2013
Recommended Dosage: Children (2.3)	10/2013
Warnings and Precautions	
Hepatic Injury (5.1)	10/2013
Serious Skin Reactions (5.2)	10/2013
Risk of Medication Errors (5.3)	10/2013

INDICATIONS AND USAGE

OFIRMEV (acetaminophen) injection is indicated for the

- Management of mild to moderate pain (1)
- Management of moderate to severe pain with adjunctive opioid analgesics (1)
- Reduction of fever (1)

DOSAGE AND ADMINISTRATION

- OFIRMEV may be given as a single or repeated dose. (2.1)
- OFIRMEV should be administered only as a 15-minute intravenous infusion. (2.4)

Adults and Adolescents Weighing 50 kg and Over:

- 1000 mg every 6 hours or 650 mg every 4 hours to a maximum of 4000 mg per day. Minimum dosing interval of 4 hours. (2.2)

Adults and Adolescents Weighing Under 50 kg:

- 15 mg/kg every 6 hours or 12.5 mg/kg every 4 hours to a maximum of 75 mg/kg per day. Minimum dosing interval of 4 hours. (2.2)

Children:

- Children 2 to 12 years of age: 15 mg/kg every 6 hours or 12.5 mg/kg every 4 hours to a maximum of 75 mg/kg per day. Minimum dosing interval of 4 hours. (2.3)

DOSAGE FORMS AND STRENGTHS

- Injection for intravenous infusion.
- Each 100 mL glass vial contains 1000 mg acetaminophen (10 mg/mL). (3)

CONTRAINDICATIONS

Acetaminophen is contraindicated:

- In patients with known hypersensitivity to acetaminophen or to any of the excipients in the IV formulation. (4)
- In patients with severe hepatic impairment or severe active liver disease. (4)

WARNINGS AND PRECAUTIONS

- Administration of acetaminophen in doses higher than recommended (by all routes of administration and from all acetaminophen-containing products including combination products) may result in hepatic injury, including the risk of liver failure and death. (5.1)
- Do not exceed the maximum recommended daily dose of acetaminophen (by all routes of administration and all acetaminophen-containing products including combination products). (5.1)
- Take care when prescribing, preparing, and administering OFIRMEV injection to avoid dosing errors which could result in accidental overdose and death. (5.3)
- Use caution when administering acetaminophen in patients with the following conditions: hepatic impairment or active hepatic disease, in cases of alcoholism, chronic malnutrition, severe hypovolemia, or severe renal impairment (creatinine clearance \leq 30 mL/min). (5.1)
- Discontinue OFIRMEV immediately at the first appearance of skin rash and if symptoms associated with allergy or hypersensitivity occur. Do not use in patients with acetaminophen allergy. (5.2, 5.4)

ADVERSE REACTIONS

The most common adverse reactions in patients treated with OFIRMEV were nausea, vomiting, headache, and insomnia in adult patients and nausea, vomiting, constipation, pruritus, agitation, and atelectasis in pediatric patients. (6.1)

To report SUSPECTED ADVERSE REACTIONS, contact Cadence Pharmaceuticals Inc. at 1-877-647-2239 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

DRUG INTERACTIONS

- Substances that induce or regulate hepatic cytochrome enzyme CYP2E1 may alter the metabolism of acetaminophen and increase its hepatotoxic potential. (7.1)
- Chronic oral acetaminophen use at a dose of 4000 mg/day has been shown to cause an increase in international normalized ratio (INR) in some patients who have been stabilized on sodium warfarin as an anticoagulant. (7.2)

USE IN SPECIFIC POPULATIONS

- Pregnancy: Category C. There are no studies of intravenous acetaminophen in pregnant women. Use only if clearly needed. (8.1)
- Nursing Mothers: Caution should be exercised when administered to a nursing woman. (8.3)
- Pediatric Use: The effectiveness of OFIRMEV for the treatment of acute pain and fever has not been studied in pediatric patients less than 2 years of age. The safety and effectiveness of OFIRMEV in pediatric patients older than 2 years is supported by evidence from adequate and well controlled studies in adults with additional safety and pharmacokinetic data for this age group. (8.4)
- Geriatric Use: No overall differences in safety or effectiveness were observed between geriatric and younger subjects. (8.5)
- Hepatic Impairment: OFIRMEV is contraindicated in patients with severe hepatic impairment or severe active liver disease and should be used with caution in patients with hepatic impairment or active liver disease. (4, 5.1, 8.6)
- Renal Impairment: In cases of severe renal impairment, longer dosing intervals and a reduced total daily dose of acetaminophen may be warranted. (5.1, 8.7)

Revised: 10/2013

FULL PRESCRIBING INFORMATION: CONTENTS*

WARNING: RISK OF MEDICATION ERRORS AND HEPATOTOXICITY

1 INDICATIONS AND USAGE

2 DOSAGE AND ADMINISTRATION

- 2.1 General Dosing Information
- 2.2 Recommended Dosage: Adults and Adolescents

- 2.3 Recommended Dosage: Children

- 2.4 Instructions for Intravenous Administration

3 DOSAGE FORMS AND STRENGTHS

4 CONTRAINDICATIONS

5 WARNINGS AND PRECAUTIONS

- 5.1 Hepatic Injury

- 5.2 Serious Skin Reactions

- 5.3 Risk of Medication Errors

- 5.4 Allergy and Hypersensitivity

6 ADVERSE REACTIONS

- 6.1 Clinical Trial Experience

7 DRUG INTERACTIONS

- 7.1 Effects of other Substances on Acetaminophen

- 7.2 Anticoagulants

8 USE IN SPECIFIC POPULATIONS

- 8.1 Pregnancy

8.2 Labor and Delivery

8.3 Nursing Mothers

8.4 Pediatric Use

8.5 Geriatric Use

8.6 Patients with Hepatic Impairment

8.7 Patients with Renal Impairment

10 OVERDOSAGE

11 DESCRIPTION

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

12.2 Pharmacodynamics

12.3 Pharmacokinetics

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

14 CLINICAL STUDIES

14.1 Adult Acute Pain

14.2 Adult Fever

14.3 Pediatric Acute Pain and Fever

16 HOW SUPPLIED/STORAGE AND HANDLING

*Sections or subsections omitted from the full prescribing information are not listed.

FULL PRESCRIBING INFORMATION**WARNING: Risk of Medication Errors and Hepatotoxicity**

Take care when prescribing, preparing, and administering OFIRMEV Injection to avoid dosing errors which could result in accidental overdose and death. In particular, be careful to ensure that:

- the dose in milligrams (mg) and milliliters (mL) is not confused;
- the dosing is based on weight for patients under 50 kg;
- infusion pumps are properly programmed; and
- the total daily dose of acetaminophen from all sources does not exceed maximum daily limits.

OFIRMEV contains acetaminophen. Acetaminophen has been associated with cases of acute liver failure, at times resulting in liver transplant and death. Most of the cases of liver injury are associated with the use of acetaminophen at doses that exceed the maximum daily limits, and often involve more than one acetaminophen-containing product (see **WARNINGS**).

1 INDICATIONS AND USAGE

OFIRMEV[®] (acetaminophen) injection is indicated for

- the management of mild to moderate pain
- the management of moderate to severe pain with adjunctive opioid analgesics
- the reduction of fever.

2 DOSAGE AND ADMINISTRATION**2.1 General Dosing Information**

OFIRMEV may be given as a single or repeated dose for the treatment of acute pain or fever. No dose adjustment is required when converting between oral acetaminophen and OFIRMEV dosing in adults and adolescents who weigh 50 kg and above. Calculated maximum daily dose of acetaminophen is based on all routes of administration (i.e., intravenous, oral, and rectal) and all products containing acetaminophen.

Exceeding the maximum mg/kg daily dose of acetaminophen as described in Tables 1 and 2 may result in hepatic injury, including the risk of liver failure and death. To avoid the risk of overdose, ensure that the total amount of acetaminophen from all routes and from all sources does not exceed the maximum recommended dose.

2.2 Recommended Dosage: Adults and Adolescents

Adults and adolescents weighing 50 kg and over: the recommended dosage of OFIRMEV is 1000 mg every 6 hours or 650 mg every 4 hours, with a maximum single dose of OFIRMEV of 1000 mg, a minimum dosing interval of 4 hours, and a maximum daily dose of acetaminophen of 4000 mg per day (includes all routes of administration and all acetaminophen-containing products including combination products).

Adults and adolescents weighing under 50 kg: the recommended dosage of OFIRMEV is 15 mg/kg every 6 hours or 12.5 mg/kg every 4 hours, with a maximum single dose of OFIRMEV of 15 mg/kg, a minimum dosing interval of 4 hours, and a maximum daily dose of acetaminophen of 75 mg/kg per day (includes all routes of administration and all acetaminophen-containing products including combination products).

Table 1. Dosing for Adults and Adolescents

Age group	Dose given every 4 hours	Dose given every 6 hours	Maximum single dose	Maximum total daily dose of acetaminophen (by all routes)
Adults and adolescents (13 years and older) weighing ≥ 50 kg	650 mg	1000 mg	1000 mg	4000 mg in 24 hours
Adults and adolescents (13 years and older) weighing < 50 kg	12.5 mg/kg	15 mg/kg	15 mg/kg (up to 750 mg)	75 mg/kg in 24 hours (up to 3750 mg)

2.3 Recommended Dosage: Children

Children 2 to 12 years of age: the recommended dosage of OFIRMEV is 15 mg/kg every 6 hours or 12.5 mg/kg every 4 hours, with a maximum single dose of OFIRMEV of 15 mg/kg, a minimum dosing interval of 4 hours, and a maximum daily dose of acetaminophen of 75 mg/kg per day.

Table 2. Dosing for Children

Age group	Dose given every 4 hours	Dose given every 6 hours	Maximum single dose	Maximum total daily dose of acetaminophen (by all routes)
Children 2 to 12 years of age	12.5 mg/kg	15 mg/kg	15 mg/kg (up to 750 mg)	75 mg/kg in 24 hours (up to 3750 mg)

2.4 Instructions for Intravenous Administration

For adult and adolescent patients weighing ≥ 50 kg requiring 1000 mg doses of OFIRMEV, administer the dose by inserting a vented intravenous set through the septum of the 100 mL vial. OFIRMEV may be administered without further dilution. Examine the vial contents before dose preparation or administering. DO NOT USE if particulate matter or discoloration is observed. Administer the contents of the vial intravenously over 15-minutes. Use aseptic technique when preparing OFIRMEV for intravenous infusion. Do not add other medications to the OFIRMEV vial or infusion device.

For doses less than 1000 mg, the appropriate dose must be withdrawn from the vial and placed into a separate container prior to administration. Using aseptic technique, withdraw the appropriate dose (650 mg or weight-based) from an intact sealed OFIRMEV vial and place the measured dose in a separate empty, sterile container (e.g. glass bottle, plastic intravenous container, or syringe) for intravenous infusion to avoid the inadvertent delivery and administration of the total volume of the commercially available container. The entire 100 mL vial of OFIRMEV is not intended for use in patients weighing less than 50 kg. OFIRMEV is a single-use vial and the unused portion must be discarded.

Place small volume pediatric doses up to 60 mL in volume in a syringe and administer over 15 minutes using a syringe pump.

Monitor the end of the infusion in order to prevent the possibility of an air embolism, especially in cases where the OFIRMEV infusion is the primary infusion.

Once the vacuum seal of the glass vial has been penetrated, or the contents transferred to another container, administer the dose of OFIRMEV within 6 hours.

Do not add other medications to the OFIRMEV solution. Diazepam and chlorpromazine hydrochloride are physically incompatible with OFIRMEV, therefore do not administer simultaneously.

3 DOSAGE FORMS AND STRENGTHS

OFIRMEV is a sterile, clear, colorless, non pyrogenic, preservative free, isotonic formulation of acetaminophen intended for intravenous infusion. Each 100 mL glass vial contains 1000 mg acetaminophen (10 mg/mL).

4 CONTRAINDICATIONS

Acetaminophen is contraindicated:

- in patients with known hypersensitivity to acetaminophen or to any of the excipients in the intravenous formulation.
- in patients with severe hepatic impairment or severe active liver disease [*see WARNINGS AND PRECAUTIONS (5.1)*].

5 WARNINGS AND PRECAUTIONS

5.1 Hepatic Injury

Administration of acetaminophen in doses higher than recommended may result in hepatic injury, including the risk of liver failure and death [*see OVERDOSAGE (10)*]. Do not exceed the maximum recommended daily dose of acetaminophen [*see DOSAGE AND ADMINISTRATION (2)*]. The maximum recommended daily dose of acetaminophen includes all routes of acetaminophen administration and all acetaminophen-containing products administered, including combination products.

Use caution when administering acetaminophen in patients with the following conditions: hepatic impairment or active hepatic disease, alcoholism, chronic malnutrition, severe hypovolemia (e.g., due to dehydration or blood loss), or severe renal impairment (creatinine clearance \leq 30 mL/min) [*see USE IN SPECIFIC POPULATIONS (8.6, 8.7)*].

5.2 Serious Skin Reactions

Rarely, acetaminophen may cause serious skin reactions such as acute generalized exanthematous pustulosis (AGEP), Stevens-Johnson Syndrome (SJS), and toxic epidermal necrolysis (TEN), which can be fatal. Patients should be informed about the signs of serious skin reactions, and use of the drug should be discontinued at the first appearance of skin rash or any other sign of hypersensitivity.

5.3 Risk of Medication Errors

Take care when prescribing, preparing, and administering OFIRMEV (acetaminophen) Injection in order to avoid dosing errors which could result in accidental overdose and death. In particular, be careful to ensure that:

- the dose in milligrams (mg) and milliliters (mL) is not confused;
- the dosing is based on weight for patients under 50 kg;
- infusion pumps are properly programmed; and
- the total daily dose of acetaminophen from all sources does not exceed maximum daily limits [*see DOSAGE AND ADMINISTRATION (2)*].

5.4 Allergy and Hypersensitivity

There have been post-marketing reports of hypersensitivity and anaphylaxis associated with the use of acetaminophen. Clinical signs included swelling of the face, mouth, and throat, respiratory distress, urticaria, rash, and pruritus. There were infrequent reports of life-threatening anaphylaxis requiring emergent medical attention. Discontinue OFIRMEV immediately if symptoms associated with allergy or hypersensitivity occur. Do not use OFIRMEV in patients with acetaminophen allergy.

6 ADVERSE REACTIONS

The following serious adverse reactions are discussed elsewhere in the labeling:

- Hepatic Injury [*see WARNINGS AND PRECAUTIONS (5.1)*]
- Serious Skin Reactions [*see WARNINGS AND PRECAUTIONS (5.2)*]
- Allergy and Hypersensitivity [*see WARNINGS AND PRECAUTIONS (5.4)*]

6.1 Clinical Trial Experience

Because clinical trials are conducted under widely varying conditions, adverse reaction rates observed cannot be directly compared to rates in other clinical trials and may not reflect the rates observed in practice.

Adult Population

A total of 1020 adult patients have received OFIRMEV in clinical trials, including 37.3% (n=380) who received 5 or more doses, and 17.0% (n=173) who received more than 10 doses. Most patients were treated with OFIRMEV 1000 mg every 6 hours. A total of 13.1% (n=134) received OFIRMEV 650 mg every 4 hours.

All adverse reactions that occurred in adult patients treated with either OFIRMEV or placebo in repeated dose, placebo-controlled clinical trials at an incidence \geq 3% and at a greater frequency than placebo are listed in Table 3. The most common adverse events in adult patients treated with OFIRMEV (incidence \geq 5% and greater than placebo) were nausea, vomiting, headache, and insomnia.

Table 3. Treatment-Emergent Adverse Reactions Occurring \geq 3% in OFIRMEV-treated Patients and at a greater frequency than Placebo in Placebo-Controlled, Repeated Dose Studies

System Organ Class – Preferred Term	OFIRMEV (N=402) n (%)	Placebo (N=379) n (%)
Gastrointestinal Disorders		
Nausea	138 (34)	119 (31)
Vomiting	62 (15)	42 (11)
General Disorders and Administration Site Conditions		
Pyrexia*	22 (5)	52 (14)
Nervous System Disorders		
Headache	39 (10)	33 (9)
Psychiatric Disorders		
Insomnia	30 (7)	21 (5)

* Pyrexia adverse reaction frequency data is included in order to alert healthcare practitioners that the antipyretic effects of OFIRMEV may mask fever.

Other Adverse Reactions Observed During Clinical Studies of OFIRMEV in Adults

The following additional treatment-emergent adverse reactions were reported by adult subjects treated with OFIRMEV in all clinical trials (n=1020) that occurred with an incidence of at least 1% and at a frequency greater than placebo (n=525).

Blood and lymphatic system disorders: anemia

General disorders and administration site conditions: fatigue, infusion site pain, edema peripheral

Investigations: aspartate aminotransferase increased, breath sounds abnormal

Metabolism and nutrition disorders: hypokalemia

Musculoskeletal and connective tissue disorders: muscle spasms, trismus

Psychiatric disorders: anxiety

Respiratory, thoracic and mediastinal disorders: dyspnea

Vascular disorders: hypertension, hypotension

Pediatric population

A total of 355 pediatric patients (47 neonates, 64 infants, 171 children, and 73 adolescents) have received OFIRMEV in active-controlled (n=250) and open-label clinical trials (n=225), including 59.7% (n=212) who received 5 or more doses and 43.1% (n=153) who received more than 10 doses. Pediatric patients received OFIRMEV doses up to 15 mg/kg on an every 4 hours, every 6 hours, or every 8 hours schedule. The maximum exposure was 7.7, 6.4, 6.8, and 7.1 days in neonates, infants, children, and adolescents, respectively.

The most common adverse events (incidence \geq 5%) in pediatric patients treated with OFIRMEV were nausea, vomiting, constipation, pruritus, agitation, and atelectasis.

Other Adverse Reactions Observed During Clinical Studies of OFIRMEV in Pediatrics

The following additional treatment-emergent adverse reactions were reported by pediatric subjects treated with OFIRMEV (n=355) that occurred with an incidence of at least 1%.

Blood and lymphatic system disorders: anemia

Cardiac disorders: tachycardia

Gastrointestinal disorders: abdominal pain, diarrhea

General disorders and administration site conditions: injection site pain, edema peripheral, pyrexia

Investigations: hepatic enzyme increase

Metabolism and nutrition disorders: hypoalbuminemia, hypokalemia, hypomagnesemia, hypophosphatemia, hypervolemia

Musculoskeletal and connective tissue disorders: muscle spasm, pain in extremity

Nervous system disorders: headache

Psychiatric disorders: insomnia

Renal and urinary disorders: oliguria

Respiratory, thoracic and mediastinal disorders: pulmonary edema, hypoxia, pleural effusion, stridor, wheezing

Skin and subcutaneous tissue disorders: periorbital edema, rash

Vascular disorders: hypertension, hypotension

7 DRUG INTERACTIONS

7.1 Effects of other Substances on Acetaminophen

Substances that induce or regulate hepatic cytochrome enzyme CYP2E1 may alter the metabolism of acetaminophen and increase its hepatotoxic potential. The clinical consequences of these effects have not been established. Effects of ethanol are complex, because excessive alcohol usage can induce hepatic cytochromes, but ethanol also acts as a competitive inhibitor of the metabolism of acetaminophen.

7.2 Anticoagulants

Chronic oral acetaminophen use at a dose of 4000 mg/day has been shown to cause an increase in international normalized ratio (INR) in some patients who have been stabilized on sodium warfarin as an anticoagulant. As no studies have been performed evaluating the short-term use of OFIRMEV in patients on oral anticoagulants, more frequent assessment of INR may be appropriate in such circumstances.

8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

Pregnancy Category C. There are no studies of intravenous acetaminophen in pregnant women; however, epidemiological data on oral acetaminophen use in pregnant women show no increased risk of major congenital malformations. Animal reproduction studies have not been conducted with IV acetaminophen, and it is not known whether OFIRMEV can cause fetal harm when administered to a pregnant woman. OFIRMEV should be given to a pregnant woman only if clearly needed.

The results from a large population-based prospective cohort, including data from 26,424 women with live born singletons who were exposed to oral acetaminophen during the first trimester, indicate no increased risk for congenital malformations, compared to a control group of unexposed children. The rate of congenital malformations (4.3%) was similar to the rate in the general population. A population-based, case-control study from the National Birth Defects Prevention Study showed that 11,610 children with prenatal exposure to acetaminophen during the first trimester had no increased risk of major birth defects compared to 4,500 children in the control group. Other epidemiological data showed similar results.

While animal reproduction studies have not been conducted with intravenous acetaminophen, studies in pregnant rats that received oral acetaminophen during organogenesis at doses up to 0.85 times the maximum human daily dose (MHDD = 4 grams/day, based on a body surface area comparison) showed evidence of

fetotoxicity (reduced fetal weight and length) and a dose-related increase in bone variations (reduced ossification and rudimentary rib changes). Offspring had no evidence of external, visceral, or skeletal malformations. When pregnant rats received oral acetaminophen throughout gestation at doses of 1.2-times the MHDD (based on a body surface area comparison), areas of necrosis occurred in both the liver and kidney of pregnant rats and fetuses. These effects did not occur in animals that received oral acetaminophen at doses 0.3-times the MHDD, based on a body surface area comparison.

In a continuous breeding study, pregnant mice received 0.25, 0.5, or 1.0% acetaminophen via the diet (357, 715, or 1430 mg/kg/day). These doses are approximately 0.43, 0.87, and 1.7 times the MHDD, respectively, based on a body surface area comparison. A dose-related reduction in body weights of fourth and fifth litter offspring of the treated mating pair occurred during lactation and post-weaning at all doses. Animals in the high dose group had a reduced number of litters per mating pair, male offspring with an increased percentage of abnormal sperm, and reduced birth weights in the next generation pups.

8.2 Labor and Delivery

There are no adequate and well-controlled studies with OFIRMEV during labor and delivery; therefore, it should be used in such settings only after a careful benefit-risk assessment.

8.3 Nursing Mothers

While studies with OFIRMEV have not been conducted, acetaminophen is secreted in human milk in small quantities after oral administration. Based on data from more than 15 nursing mothers, the calculated infant daily dose of acetaminophen is approximately 1 – 2% of the maternal dose. There is one well-documented report of a rash in a breast-fed infant that resolved when the mother stopped acetaminophen use and recurred when she resumed acetaminophen use. Caution should be exercised when OFIRMEV is administered to a nursing woman.

8.4 Pediatric Use

The safety and effectiveness of OFIRMEV for the treatment of acute pain and fever in pediatric patients ages 2 years and older is supported by evidence from adequate and well-controlled studies of OFIRMEV in adults. Additional safety and pharmacokinetic data were collected in 355 patients across the full pediatric age strata, from premature neonates (≥ 32 weeks post menstrual age) to adolescents. The effectiveness of OFIRMEV for the treatment of acute pain and fever has not been studied in pediatric patients < 2 years of age. [see DOSAGE AND ADMINISTRATION - Recommended Dosage: Children (2.3) and PHARMACOKINETICS (12.3)].

8.5 Geriatric Use

Of the total number of subjects in clinical studies of OFIRMEV, 15% were age 65 and over, while 5% were age 75 and over. No overall differences in safety or effectiveness were observed between these subjects and younger subjects, and other reported clinical experience has not identified differences in responses between the elderly and younger patients, but greater sensitivity of some older individuals cannot be ruled out.

8.6 Patients with Hepatic Impairment

Acetaminophen is contraindicated in patients with severe hepatic impairment or severe active liver disease and should be used with caution in patients with hepatic impairment or active liver disease [see WARNINGS AND PRECAUTIONS (5.1), CLINICAL PHARMACOLOGY (12)]. A reduced total daily dose of acetaminophen may be warranted.

8.7 Patients with Renal Impairment

In cases of severe renal impairment (creatinine clearance ≤ 30 mL/min), longer dosing intervals and a reduced total daily dose of acetaminophen may be warranted.

10 OVERDOSAGE

Signs and Symptoms

In acute acetaminophen overdosage, dose-dependent, potentially fatal hepatic necrosis is the most serious adverse effect. Renal tubular necrosis, hypoglycemic coma, and thrombocytopenia may also occur. Plasma acetaminophen levels > 300 mcg/mL at 4 hours after oral ingestion were associated with hepatic damage in

90% of patients; minimal hepatic damage is anticipated if plasma levels at 4 hours are < 150 mcg/mL or < 37.5 mcg/mL at 12 hours after ingestion. Early symptoms following a potentially hepatotoxic overdose may include: nausea, vomiting, diaphoresis, and general malaise. Clinical and laboratory evidence of hepatic toxicity may not be apparent until 48 to 72 hours post-ingestion.

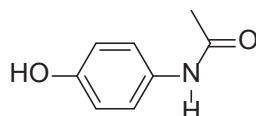
Treatment

If an acetaminophen overdose is suspected, obtain a serum acetaminophen assay as soon as possible, but no sooner than 4 hours following oral ingestion. Obtain liver function studies initially and repeat at 24-hour intervals. Administer the antidote N-acetylcysteine (NAC) as early as possible. As a guide to treatment of acute ingestion, the acetaminophen level can be plotted against time since oral ingestion on a nomogram (Rumack-Matthew). The lower toxic line on the nomogram is equivalent to 150 mcg/mL at 4 hours and 37.5 mcg/mL at 12 hours. If serum level is above the lower line, administer the entire course of NAC treatment. Withhold NAC therapy if the acetaminophen level is below the lower line.

For additional information, call a poison control center at 1-800-222-1222.

11 DESCRIPTION

Acetaminophen is a non-salicylate antipyretic and non-opioid analgesic agent. Its chemical name is N-acetyl-p-aminophenol. Acetaminophen has a molecular weight of 151.16. Its structural formula is:



OFIRMEV injection is a sterile, clear, colorless, non pyrogenic, isotonic formulation of acetaminophen intended for intravenous infusion. It has a pH of approximately 5.5 and an osmolality of approximately 290 mOsm/kg. Each 100 mL contains 1000 mg acetaminophen, USP, 3850 mg mannitol, USP, 25 mg cysteine hydrochloride, monohydrate, USP, 10.4 mg dibasic sodium phosphate, USP. pH is adjusted with hydrochloric acid and/or sodium hydroxide.

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

The precise mechanism of the analgesic and antipyretic properties of acetaminophen is not established but is thought to primarily involve central actions.

12.2 Pharmacodynamics

Acetaminophen has been shown to have analgesic and antipyretic activities in animal and human studies.

Single doses of OFIRMEV up to 3000 mg and repeated doses of 1000 mg every 6 hours for 48 hours have not been shown to cause a significant effect on platelet aggregation. Acetaminophen does not have any immediate or delayed effects on small-vessel hemostasis. Clinical studies of both healthy subjects and patients with hemophilia showed no significant changes in bleeding time after receiving multiple doses of oral acetaminophen.

12.3 Pharmacokinetics

Distribution

The pharmacokinetics of OFIRMEV have been studied in patients and healthy subjects from premature neonates up to adults 60 years old. The pharmacokinetic profile of OFIRMEV has been demonstrated to be dose proportional in adults following administration of single doses of 500, 650, and 1000 mg.

The maximum concentration (C_{max}) occurs at the end of the 15 minute intravenous infusion of OFIRMEV. Compared to the same dose of oral acetaminophen, the C_{max} following administration of OFIRMEV is up to 70% higher, while overall exposure (area under the concentration time curve [AUC]) is very similar.

Pharmacokinetic parameters of OFIRMEV (AUC, C_{max} , terminal elimination half-life [$T_{1/2}$], systemic clearance [CL], and volume of distribution at steady state [V_{ss}]) following administration of a single intravenous dose of 15 mg/kg for the pediatric population and 1000 mg in adults are summarized in Table 4.

Table 4. OFIRMEV Pharmacokinetic Parameters

Subpopulations	Mean (SD)				
	AUC ($\mu\text{g} \times \text{h/mL}$)	C_{max} ($\mu\text{g/mL}$)	$\text{T}_{\frac{1}{2}}$ (h)	CL (L/h/kg)	V_{ss} (L/kg)
Neonates	62 (11)	25 (4)	7.0 (2.7)	0.12 (0.04)	1.1 (0.2)
Infants	57 (54)	29 (24)	4.2 (2.9)	0.29 (0.15)	1.1 (0.3)
Children	38 (8)	29 (7)	3.0 (1.5)	0.34 (0.10)	1.2 (0.3)
Adolescents	41 (7)	31 (9)	2.9 (0.7)	0.29 (0.08)	1.1 (0.3)
Adults	43 (11)	28 (21)	2.4 (0.6)	0.27 (0.08)	0.8 (0.2)

The pharmacokinetic exposure of OFIRMEV observed in children and adolescents is similar to adults, but higher in neonates and infants. Dosing simulations from pharmacokinetic data in infants and neonates suggest that dose reductions of 33% in infants 1 month to < 2 years of age, and 50% in neonates up to 28 days, with a minimum dosing interval of 6 hours, will produce a pharmacokinetic exposure similar to that observed in children age 2 years and older.

At therapeutic levels, binding of acetaminophen to plasma proteins is low (ranging from 10% to 25%). Acetaminophen appears to be widely distributed throughout most body tissues except fat.

Metabolism and Excretion

Acetaminophen is primarily metabolized in the liver by first-order kinetics and involves three principal separate pathways: Conjugation with glucuronide, conjugation with sulfate, and oxidation via the cytochrome P450 enzyme pathway, primarily CYP2E1, to form a reactive intermediate metabolite (N-acetyl-p-benzoquinone imine or NAPQI). With therapeutic doses, NAPQI undergoes rapid conjugation with glutathione and is then further metabolized to form cysteine and mercapturic acid conjugates.

Acetaminophen metabolites are mainly excreted in the urine. Less than 5% is excreted in the urine as unconjugated (free) acetaminophen and more than 90% of the administered dose is excreted within 24 hours.

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

Carcinogenesis

Long-term studies in mice and rats have been completed by the National Toxicology Program to evaluate the carcinogenic potential of acetaminophen. In 2-year feeding studies, F344/N rats and B6C3F1 mice were fed a diet containing acetaminophen up to 6000 ppm. Female rats demonstrated equivocal evidence of carcinogenic activity based on increased incidences of mononuclear cell leukemia at 0.8 times the maximum human daily dose (MHDD) of 4 grams/day, based on a body surface area comparison. In contrast, there was no evidence of carcinogenic activity in male rats (0.7 times) or mice (1.2-1.4 times the MHDD, based on a body surface area comparison).

Mutagenesis

Acetaminophen was not mutagenic in the bacterial reverse mutation assay (Ames test). In contrast, acetaminophen tested positive in the in vitro mouse lymphoma assay and the in vitro chromosomal aberration assay using human lymphocytes. In the published literature, acetaminophen has been reported to be clastogenic when administered a dose of 1500 mg/kg/day to the rat model (3.6-times the MHDD, based on a body surface area comparison). In contrast, no clastogenicity was noted at a dose of 750 mg/kg/day (1.8-times the MHDD, based on a body surface area comparison), suggesting a threshold effect.

Impairment of fertility

In studies conducted by the National Toxicology Program, fertility assessments have been completed in Swiss mice via a continuous breeding study. There were no effects on fertility parameters in mice consuming up to 1.7 times the MHDD of acetaminophen, based on a body surface area comparison. Although there was no effect on sperm motility or sperm density in the epididymis, there was a significant increase in the percentage of abnormal sperm in mice consuming 1.7 times the MHDD (based on a body surface area comparison) and there

was a reduction in the number of mating pairs producing a fifth litter at this dose, suggesting the potential for cumulative toxicity with chronic administration of acetaminophen near the upper limit of daily dosing.

Published studies in rodents report that oral acetaminophen treatment of male animals at doses that are 1.2 times the MHDD and greater (based on a body surface area comparison) result in decreased testicular weights, reduced spermatogenesis, reduced fertility, and reduced implantation sites in females given the same doses. These effects appear to increase with the duration of treatment. The clinical significance of these findings is not known.

14 CLINICAL STUDIES

14.1 Adult Acute Pain

The efficacy of OFIRMEV in the treatment of acute pain in adults was evaluated in two randomized, double-blind, placebo-controlled clinical trials in patients with postoperative pain.

Pain Study 1 evaluated the analgesic efficacy of repeated doses of OFIRMEV 1000 mg vs. placebo every 6 hours for 24 hours in 101 patients with moderate to severe pain following total hip or knee replacement. OFIRMEV was statistically superior to placebo for reduction in pain intensity over 24 hours. There was an attendant decrease in opioid consumption, the clinical benefit of which was not demonstrated.

Pain Study 2 evaluated the analgesic efficacy of repeated doses of OFIRMEV 1000 mg every 6 hours or 650 mg every 4 hours for 24 hours versus placebo in the treatment of 244 patients with moderate to severe postoperative pain after abdominal laparoscopic surgery. Patients receiving OFIRMEV experienced a statistically significant greater reduction in pain intensity over 24 hours compared to placebo.

14.2 Adult Fever

The efficacy of OFIRMEV 1000 mg in the treatment of adult fever was evaluated in one randomized, double-blind, placebo-controlled clinical trial. The study was a 6-hour, single-dose, endotoxin-induced fever study in 60 healthy adult males. A statistically significant antipyretic effect of OFIRMEV was demonstrated through 6 hours in comparison to placebo. The mean temperature over time is shown in Figure 1.

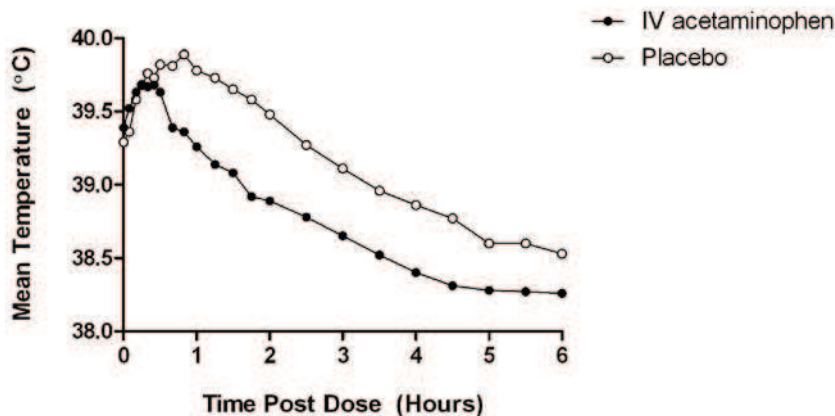


Figure 1: Mean Temperature (°C) Over Time

14.3 Pediatric Acute Pain and Fever

OFIRMEV was studied in 355 pediatric patients in two active-controlled and three open-label safety and pharmacokinetic trials [see PEDIATRIC USE (8.4)].

16 HOW SUPPLIED/STORAGE AND HANDLING

OFIRMEV is supplied in a 100 mL glass vial containing 1000 mg acetaminophen (10 mg/mL).

Carton of 24 vials, NDC 43825-102-01

OFIRMEV should be stored at 20 °C to 25 °C (68 °F to 77 °F) [See USP Controlled Room Temperature]. For single use only. The product should be used within 6 hours after opening. Do not refrigerate or freeze.

OFIRMEV (acetaminophen) Injection

Manufactured for:
Cadence Pharmaceuticals, Inc.
San Diego, CA 92130

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Label part number